



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center  
Office of Analytical Services and Quality Assurance  
701 Mapes Road  
Fort Meade, Maryland 20755-5350



**Final Analytical Report**

Site Name.....	Dimock Residential Groundwater
Sample Collection Date(s).....	01/23/12 12:30- 01/27/12 13:18
Contact.....	Rich Fetzer
Report Date.....	02/11/12 15:37
Project #.....	DAS R33907
Work Order.....	1201013

**Analyses included in this report:**

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Alcohols by EPA 8015D	SVOCs by CLP Equivalent
VOCs by CLP Equivalent (trace)	

Approved for Release

1201013 FINAL      DAS R33907      02 11 12 1538  
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OASQA Representative



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center  
Office of Analytical Services and Quality Assurance  
701 Mapes Road  
Fort Meade, Maryland 20755-5350



**Site Name: Dimock Residential Groundwater**

**Project #: DAS R33907**

Report Narrative

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1201013 FINAL

DAS R33907

02 11 12 1538

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**Site Name: Dimock Residential Groundwater**

**Project #: DAS R33907**

**Report Narrative**

The EPA Region 3 Laboratory's Quality System is NELAP accredited. The National Environmental Laboratory Accreditation Program (NELAP) is a voluntary environmental laboratory accreditation association of State and Federal agencies.

**General Notes:**

This report contains results for Volatiles (VOAs), Semivolatiles (SVOAs), and Alcohol analyses only. All other parameters identified on the chain-of-custody form are included in separate reports. Lab Sample numbers 1201013-02, -04, -06, -08, -10, -18, -19, -21 thru -24, -27, 37 thru -44 are not included in this report since these samples were designated for Metals and Mercury analyses only.

For Work Order 1201013 - **This is Report 2 of 3.**

All samples were received intact and at proper temperature.

Lab Sample number 1201013-20 was cancelled. This sample was designated for Oil & Grease analysis and shipped independently from the other analytical fractions. The sample was reassigned to Lab Sample number 1201013-33.

For samples received by the laboratory on 1/28/2012, the analyses Alcohol and Glycol for FB03 (1201013-12) were not recorded on the chain-of-custody form. A request for a Letter-to-File was submitted to the sampler on 2/9/2012.

Some samples designated for the analysis of Orthophosphorous were received at the laboratory past the established holding times. Therefore, all samples were analyzed using the Total Phosphate method and results for the analysis by the Orthophosphorous method are not included in this report. Since the Orthophosphorous method was being used as a screening method to determine the need to analyze the sample by the Total Phosphate method, results for Total Phosphate are not impacted.

Samples designated for the analysis of Oil & Grease were received in sample containers inconsistent with the type needed for the routine extraction procedure. Therefore, all samples were extracted using the manual extraction technique.

Where applicable, sample results are qualified based on the highest level concentrations of field QC contamination found in the field, equipment, or trip blanks.

**SVOAs Analysis Note:**

All samples were extracted by EPA SW-846 Method 3520C followed by analysis using EPA SW-846 Method 8270D. Refer to notes in case file for additional information regarding the analysis.

A separate calibration curve is used for two compounds, 2-methoxyethanol and 1-methylnaphthalene, with quality control requirements per the on-demand protocol.

Quantitation limit for 2,4-Dinitrophenol is qualified estimated "UJ" in sample 1201013-01 due to zero percent recovery in the low-spike quality control check. Quantitation limits for 2,4-Dinitrophenol, pentachlorophenol, and 4,6-dinitro-2-methylphenol are qualified estimated "UJ" in samples 1201013-03, -05, -07, 09, -12 thru -17, -28-36 due to low recovery in the low-spike quality control checks.

**VOA Analysis Note:**

Acrylonitrile was analyzed on-demand using CLP equivalent methodology. This analyte does not appear in the data tables or the QC summary and all data for this compound is summarized here. Acrylonitrile was not detected in any of the samples above a quantitation limit of 2 ug/L. A four point curve was analyzed (2, 5, 10 and 20 ug/L). The samples were preserved to a pH<2 with HCl. A low level second source blank spike analyzed at a concentration of 2 ug/L had a recovery of 112%. A mid level second source blank spike



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**Report Narrative**

analyzed at a concentration of 5 ug/L had a recovery of 102%. A duplicate second source blank spike at 5 ug/L had 205% recovery. Due to this high recovery, duplicate blank spikes from the primary source were analyzed. Recoveries for these spikes were 110% and 157%.

2-Chloroethylvinyl ether is not included in the analysis. 2-chloroethylvinyl ether breaks down in acidified samples.

Matrix spike/matrix spike duplicate analysis could not be completed due to insufficient sample volume. A single matrix spike was performed for samples 1201013-14 and 1201013-33.

The acetone result for sample 1201013-36 is qualified with a "K" due to an interference from isopropanol.

**Alcohols Analysis Note:**

None.

**REPORT 2 of 3**



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ANALYTICAL REPORT FOR SAMPLES

Table with 5 columns: Station ID, Laboratory ID, Matrix, Date Sampled, Date Received. Contains 30 rows of sample data.



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907



USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No: L

Chain of Custody Record
Date Shipped: 1/24/2012
Carrier Name: FedEx
Airbill: 841769084246
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr
Relinquished By: [Signature]
Received By: [Signature]

Table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNAROUND, TAG No./ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Row 1: FB01, Aqueous/ Bryan Berna, L/G, Anions (7), GC, Alcohol (7), Glycols (5/PR), Metals (5/PR), OilGrease (7), 15 (-NA- / 40mlGlassVial), 16 (-NA- / 40mlGlassVial), 17 (-NA- / 500mlHDPE), 18 (-NA- / 40mlGlassVial), 19 (HNO3 / 500mlHDPE), 21 (H2SO4 / 1000ml (WM)Amber) (6), FB01, S: 1/23/2012 14:42, 1201013-01

Kevin Martin
Kevin Martin 1/25/12
10:40

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s): [Signature]
Cooler Temperature Upon Receipt: 5°C
Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact? Shipment Iced?

TR Number: 3-043013577-012412-0012

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
Send Copy to: Sample Management Office, Attn: Heather Bauer, CSC, 15000 Conference Center Dr., Chantilly, VA 20151-3819; Phone 703/818-4200; Fax 703/818-4602

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701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907



USEPA Contract Laboratory Program
Generic Chain of Custody

Reference Case

Client No: CT5865
SDG No:

L

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Form containing shipping details (Date Shipped: 1/24/2012, Carrier Name: FedEx, Airbill: 841769084246) and Chain of Custody Record table with columns for Relinquished By, Received By, Date/Time.

Table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNAROUND, TAG No/ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes rows for FB01 and FB01-F.

Handwritten signature: Kevin Martin, 1/27/12 10:40

Form with fields: Shipment for Case Complete?, Sample(s) to be used for laboratory QC, Additional Sampler Signature(s), Cooler Temperature Upon Receipt: 5°C, Chain of Custody Seal Number, Analysis Key, Concentration, Type/Designate, Custody Seal Intact?, Shipment Iced?.

TR Number: 3-043013577-012412-0013

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program Generic Chain of Custody. This Page Left Blank. Reference Case: Client No: CT5865, SDG No: L. For Lab Use Only: Lab Contract No, Unit Price, Transfer To, Lab Contract No, Unit Price.

Table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No./PRESERVATIVE/BOTTLES, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Row 1: HW19, Drinking Water/Mike Ferrier, L/G, Anions (7), GC\_Alcohol (7), Glycols (5/PR), Metals (5/PR), Oil/Grease (7), 31 (-NA- / 40mlGlassVial), 32 (-NA- / 40mlGlassVial), 33 (-NA- / 500mlHDPE), 34 (-NA- / 40mlGlassVial), 35 (HNO3 / 500mlHDPE), 37 (H2SO4 / 1000ml (WM)Amber) (6), HW19, S: 1/23/2012 17:47, 1201013-03

Handwritten signatures: Kevin Martin, Kim Martin 1/25/12 10:40

Shipment for Case Complete?, Sample(s) to be used for laboratory QC, Additional Sampler Signature(s), Cooler Temperature Upon Receipt, Chain of Custody Seal Number, Analysis Key, Concentration, Type/Designate, Custody Seal Intact?, Shipment Iced?, Anions = 07-Anions\_Method 300.0, GC\_Alcohol = 07-Alcohols, Glycols = 07-Glycols+2-Butoxyethanol+EGME, Metals = 07-Metals, Oil/Grease = C7-Oil & Grease, HEM

TR Number: 3-043013577-012412-0014

LABORATORY COPY

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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No:

Chain of Custody Record
Date Shipped:
Carrier Name: FedEx
Airbill:
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr
Relinquished By: B. Bunnell 1/25/12 10:00
Received By: Kevin Martin 1/26/12 11:50

Table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No./PRESERVATIVE/Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes rows for HW19, HW19-F, HW19-P, and HW19-PF.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s):
Cooler Temperature Upon Receipt: 2°C
Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact?
Shipment Iced?

TR Number: 3-043013577-012512-0001

LABORATORY COPY

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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No:
For Lab Use Only
Lab Contract No:
Unit Price:
Transfer To:
Lab Contract No:
Unit Price:

Chain of Custody Record
Relinquished By (Date / Time)
Received By (Date / Time)
2
3
4

Table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNAROUND, TAG No./ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes rows for FB02 and FB02-F.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s):
Cooler Temperature Upon Receipt:
Chain of Custody Seal Number:
Analysis Key:
Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact?
Shipment Iced?

TR Number: 3-043013577-012612-0003

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Site Name: Dimock Residential Groundwater

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EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No: L

Chain of Custody Record
Relinquished By (Date / Time) Received By (Date / Time)
For Lab Use Only
Lab Contract No:
Unit Price:
Transfer To:
Lab Contract No:
Unit Price:

Table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNAROUND, TAG No/ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes rows for HW04 and HW04-F.

Shipment for Case Complete? Sample(s) to be used for laboratory QC: Additional Sampler Signature(s): Cooler Temperature Upon Receipt: Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High Type/Designate: Composite = C, Grab = G Custody Seal Intact? Shipment Iced?
Anions = 07-Anions\_Method 300.0, F-Metals = 07-Metals Dissolved, Metals = 07-Metals, Oil/Grease = 07-Oil & Grease, HEM, SVOC+TIC = 07-Semivolatiles (TCL+TICs+EGME+1-Methy, TDS = 07-Solids\_Total Dissolved, TSS = 07-Solids\_Total Suspended, WetChem = 07-Phosphorous, NO3- NO2- Tot-N

TR Number: 3-043013577-012512-0008

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
Send Copy to: Sample Management Office, Attn: Heather Bauer, CSC, 15000 Conference Center Dr., Chantilly, VA 20151-3819; Phone 703/818-4200; Fax 703/818-4602
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701 Mapes Road
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No:
For Lab Use Only
Lab Contract No:
Unit Price:
Transfer To:
Lab Contract No:
Unit Price:

Chain of Custody Record
Relinquished By (Date / Time) Received By (Date / Time)
1 B. Bowers 1/24/12 11:00 John D. Wray 1/27/12 11:59

Table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNAROUND, TAG No./ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Rows include FB02 and HW04.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s):
Cooler Temperature Upon Receipt:
Chain of Custody Seal Number:
Analysis Key:
Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact? Shipment Iced?

TR Number: 3-043013577-012612-0004

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Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No:
L

Chain of Custody Record
Date Shipped: 1/26/2012
Carrier Name: FedEx
Airbill: 8417 6908 4316
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr
Relinquished By: Brian Burris 1/26/12 08:00
Received By: John D. Cooney 1/27/12 11:59

Table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNAROUND, TAG No./ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Rows include FB01, FB02, HW04, HW19, HW19-P, TB01.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s): Bryan Berna, Tom Sedlacek, Mike Ferrier
Cooler Temperature Upon Receipt: 4.0°C
Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact? Shipment Iced?

TR Number: 3-043013577-012612-0002

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701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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USEPA Chain of Custody Record No: 3-012712-135301-0004
Date Shipped: 1/27/2012 Site #: A3TA Lab: EPA R3 Laboratory
Carrier Name: FedEx Case Complete: False Lab Contact:
Airbill No: 7979 9545 8780 Lab Phone: 410.305.3032

Table with 7 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Row 1: HW06, Drinking Water/Bryan Berna, Grab, GC\_Alcohol(7), GC\_Alcohol(7), Glycols, SVOC+TIC, SVOC+TIC, VOC+TIC(7), VOC+TIC(7), VOC+TIC(7), VOC+TIC(7), 511 (-NA- / 40mlGlassVial), 512 (-NA- / 40mlGlassVial), 514 (-NA- / 40mlGlassVial), 518 (-NA- / 1000mlAmber), 519 (-NA- / 1000mlAmber), 522 (HCl / 40mlGlassVial), 523 (HCl / 40mlGlassVial), 524 (HCl / 40mlGlassVial), 525 (HCl / 40mlGlassVial) (9), HW06, 01/26/2012 15:30, /20/DIS-17

Special Instructions: 7°C Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #
Analysis Key: GC\_Alcohol=07-Alcohols, Anions=07-Anions, Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, OilGrease=07-Oil & Grease, HEM, WetChem=07-Phosphorous, NO3-, NO2-, Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen), TDS=07-Solids, Total Dissolved, TSS=07-Solids, Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Row 1: 9, Del By, 1/27/12, K. M., 1/28/12, 14:11



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Site Name: Dimock Residential Groundwater

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USEPA Contract No. 68-11-6-002V
Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill No: 7979 9545 8780

CHAIN OF CUSTODY RECORD
Site #: A3TA
Case Complete: False

No: 3-012712-135301-0004
Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 7 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains two rows of sample data.

Special Instructions: 7°C Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains one row of handwritten data.



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill: 7931 6427 9021
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr, 701 Mapes Road, Ft. Meade MD 20755-5350

Chain of Custody Record table with columns: Relinquished By, Date/Time, Received By, Date/Time. Includes handwritten signatures and dates.

Reference Case
Client No: CT5865
SDG No: L

For Lab Use Only
Lab Contract No:
Unit Price:
Transfer To:
Lab Contract No:
Unit Price:

Main data table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No/PRESERVATIVE/BOTTLES, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY. Includes handwritten sample numbers like 1201013-12.

Shipment for Case Complete? Sample(s) to be used for laboratory QC: Additional Sampler Signature(s): Cooler Temperature Upon Receipt: Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High Type/Designate: Composite = C, Grab = G Custody Seal Intact? Shipment Iced?

TR Number: 3-043013577-012712-0004

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701 Mapes Road
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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USEPA SLI Generic Chain of Custody Record
Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill No: 7979 9545 6941
Site #: A3TA
Case Complete: False
No: 3-012712-143507-0005
Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains rows for samples HW06, HW06-F, and HW12.

Special Instructions: 7°C Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains one entry for item 8.



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No: L

Chain of Custody Record
Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill: 8417 6908 4350
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr, 701 Mapes Road, Ft. Meade MD 20755-5350
Relinquished By: [Signature] (Date / Time)
Received By: [Signature] (Date / Time)

Table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No./PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes rows for FB03, FB03F, and HW02 with detailed analysis and collection notes.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s): [Signature] for Mike Ferrier
Cooler Temperature Upon Receipt: 8°C
Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact?
Shipment Iced?

TR Number: 3-043013577-012712-0008

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
Send Copy to: Sample Management Office, Attn: Heather Bauer, CSC, 15000 Conference Center Dr., Chantilly, VA 20151-3819; Phone 703/818-4200; Fax 703/818-4602



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No: L

Chain of Custody Record table with columns: Date Shipped, Carrier Name, Airbill, Shipped to, Relinquished By, Received By, Sampler Signature, For Lab Use Only (Lab Contract No, Unit Price, Transfer To)

Main data table with columns: SAMPLE No., MATRIX/ SAMPLER, CONC/ TYPE, ANALYSIS/ TURNOVER, TAG No./ PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt

Shipment for Case Complete? table with fields: Sample(s) to be used for laboratory QC, Additional Sampler Signature(s), Cooler Temperature Upon Receipt, Chain of Custody Seal Number, Analysis Key, Concentration, Type/Designate, Custody Seal Intact?, Shipment Iced?

TR Number: 3-043013577-012712-0005

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

Reference Case
Client No: CT5865
SDG No: L

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Chain of Custody Record table with columns: Date Shipped, Carrier Name, Airbill, Shipped to, Relinquished By, TAG No, STATION LOCATION, ANALYSIS, PRESERVATIVE, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt

Main data table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No/PRESERVATIVE/Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt

Handwritten notes: HW02z Drinking Water/ Mike Ferrer L/G GC-Alcohol (7) 336 (N/A / 40 ml vial) 337 (N/A / 40 ml vial) (2) HW02z 1/25/2012 1259 (S) 1201013-14 1/27/12

Shipment for Case Complete? Sample(s) to be used for laboratory QC: Additional Sampler Signature(s): Cooler Temperature Upon Receipt: Chain of Custody Seal Number: Analysis Key: Concentration: L = Low, M = Low/Medium, H = High Type/Designate: Composite = C, Grab = G Custody Seal Intact? Shipment Iced? Anions = 07-Anions, Method 300.0, F-Metals = 07-Metals Dissolved, GC\_Alcohol = 07-Alcohols, Metals = 07-Metals, Oil/Grease = 07-Oil & Grease, HEM, TDS = 07-Solids, Total Dissolved, TSS = 07-Solids Total Suspended, VOC+TIC = 07-Volatiles+Acrylonitrile (TCI+TICs)

TR Number: 3-043013577-012712-0004

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs. Send Copy to: Sample Management Office, Attn: Heather Bauer, CSC, 15000 Conference Center Dr., Chantilly, VA 20151-3819; Phone 703/818-4200; Fax 703/818-4602

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody
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Reference Case
Client No: CT5865
SDG No:
L

Chain of Custody Record
Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill: 7979 9357 9777
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr
Relinquished By: [Signature] (Date/Time)
Received By: [Signature] (Date/Time)
For Lab Use Only
Lab Contract No:
Unit Price:
Transfer To:
Lab Contract No:
Unit Price:

Table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No./PRESERVATIVE/BOTTLES, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes rows for HW02, HW02z, and HW08a.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s):
Cooler Temperature Upon Receipt: 6°C
Chain of Custody Seal Number:
Analysis Key:
Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact?
Shipment Iced?

TR Number: 3-043013577-012712-0001

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
Send Copy to: Sample Management Office, Attn: Heather Bauer, CSC, 15000 Conference Center Dr., Chantilly, VA 20151-3819; Phone 703/818-4200; Fax 703/818-4602

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

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Reference Case
Client No: CT5865
SDG No: L

Chain of Custody Record
Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill: 7979 9552 9640
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr
Relinquished By: [Signature] 1/27/12 14:30
Received By: [Signature] 1/28/12 14:11

Table with columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No./PRESERVATIVE/ Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY. Includes rows for HW08a and HW08a-F.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC: HW08a, HW08a-F
Additional Sampler Signature(s):
Cooler Temperature Upon Receipt: 7C
Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact? Shipment Iced?

TR Number: 3-043013577-012712-0006
LABORATORY COPY
PR provides preliminary results. Requests for preliminary results will increase analytical costs.
Send Copy to: Sample Management Office, Attn: Heather Bauer, CSC, 15000 Conference Center Dr., Chantilly, VA 20151-3819; Phone 703/618-4200; Fax: 703/618-4672



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

EPA USEPA Contract Laboratory Program
Generic Chain of Custody

Reference Case
Client No: CT5865
SDG No: L

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Chain of Custody Record
Date Shipped: 1/27/2012
Carrier Name: FedEx
Airbill: 7979 9357 9777
Shipped to: US EPA, Region 3, OASQA, Env. Science Ctr, 701 Mapes Road, Ft. Meade MD 20755-5350
Relinquished By: [Signature] 1/27/12 @ 0930
Received By: [Signature] 1/28/12 14:11

Table with 8 columns: SAMPLE No., MATRIX/SAMPLER, CONC/TYPE, ANALYSIS/TURNAROUND, TAG No./PRESERVATIVE/Bottles, STATION LOCATION, SAMPLE COLLECT DATE/TIME, FOR LAB USE ONLY Sample Condition On Receipt. Includes handwritten sample IDs like 1201013-23.

Shipment for Case Complete?
Sample(s) to be used for laboratory QC:
Additional Sampler Signature(s): [Signature]
Cooler Temperature Upon Receipt: 6°C
Chain of Custody Seal Number:
Analysis Key: Concentration: L = Low, M = Low/Medium, H = High
Type/Designate: Composite = C, Grab = G
Custody Seal Intact? Shipment Iced?

TR Number: 3-043013577-012712-0002

LABORATORY COPY

PR provides preliminary results. Requests for preliminary results will increase analytical costs.
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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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USEPA CDP General VOC (LAB COPY)

CHAIN OF CUSTODY RECORD

No: 3-012812-145544-0020

Date of Report: 1/29/12

Site #: A3TA

Lab: EPA R3 Laboratory

Carrier Name: FedEx

Case Complete: False

Lab Contact:

Airbill No: 7979 9818 0407

Lab Phone: 410.305.3032

Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains rows for samples FB04, FB04-F, and HW14.

Special Instructions: 3°C temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Includes handwritten entries for 2019 and DB 1/29/12.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

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701 Mapes Road
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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CHAIN OF CUSTODY RECORD

No: 3-012812-111141-0012

OS: PA C P Gen... (LAB COPY)
Date Shipped: 1/28/2012
Carrier Name: FedEx
Airbill No: 7979 9807 0285

Site #: A3TA
Case Complete: False

Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 7 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains rows for samples FB05, FB05-F, and HW24.

Special Instructions: 2°C temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #
Analysis Key: GC\_Alcohol=07-Alcohols, Anions=07-Anions, Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, OilGrease=07-Oil & Grease, HEM, WetChem=07-Phosphorous, NO3-, NO2-, Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen, TDS=07-Solids, Total Dissolved, TSS=07-Solids, Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains one entry for item 19.



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Region 3 Environmental Science Center
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701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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CHAIN OF CUSTODY RECORD

No: 3-012812-105237-0009

U.S. EPA Form 806 (1-15-2009)

Date Shipped: 1/28/2012

Site #: A3TA

Lab: EPA R3 Laboratory

Carrier Name: FedEx

Case Complete: False

Lab Contact:

Airbill No: 7979 9807 0300

Lab Phone: 410.305.3032

Table with 7 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains two rows of sample data with handwritten notes.

Special Instructions: 30c temp Blank. Shipment for Case Complete? N. Samples Transferred From Chain of Custody #.

Analysis Key: GC\_Alcohol=07-Alcohols, Anions=07-Anions, Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, WetChem=07-Phosphorous\_NO3-,NO2-,Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen, TDS=07-Solids, Total Dissolved, TSS=07-Solids Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains one row of data with handwritten entries.



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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SEPA Chain of Custody Record
Date Shipped: 1/28/2012
Carrier Name: FedEx
Airbill No: 797998070399

No: 3-012812-103308-0008
Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 7 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains 3 rows of sample data.

Special Instructions: 20c Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains 1 row of data.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

Page 1 of 1

SELECTION General (DGC LAB COPY) CHAIN OF CUSTODY RECORD
Date Shipped: 1/28/2012 Site #: A3TA

No: 3-012812-112448-0013

Carrier Name: FedEx
Airbill No: 7979 8907 0425

Case Complete: False

Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains rows for samples HW14, HW14-F, and HW14-P.

Special Instructions: 30C Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains handwritten entry for item 14.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
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701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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SE: [unclear] General [unclear] (1/28/2012)
Date Shipped: 1/28/2012
Carrier Name: FedEx
Airbill No: 7979 9807 0436
CUSTODY RECORD
Site #: A3TA
Case Complete: False

No: 3-012812-110206-0010
Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Row 1 contains data for sample TB05.

Special Instructions: no temp Blank (cooler was packed with ice)
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #
Analysis Key: GC, Alcohol=07-Alcohols, Anions=07-Anions, Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, WetChem=07-Phosphorous, NO3-, NO2-, Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen, TDS=07-Solids, Total Dissolved, TSS=07-Solids, Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Row 1 contains handwritten data.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
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Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

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USEPA Chain of Custody Form (USEPA Form 1601-1)
Date Shipped: 1/28/2012
Carrier Name: FedEx
Airbill No: 7979 9807 2063

No: 3-012812-121815-0017
Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains 3 rows of sample data.

Special Instructions: 40C Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains 1 row of data.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

Page 2 of 2

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USEPA Contract # 68-01-0001 (LAB COPY)
Date Shipped: 1/28/2012
Carrier Name: FedEx
Airbill No: 7979 9807 0285

CHAIN OF CUSTODY RECORD
Site #: A3TA
Case Complete: False

No: 3-012812-111141-0012
Lab: EPA R3 Laboratory
Lab Contact:
Lab Phone: 410.305.3032

Table with 7 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains two rows of data for samples HW24-P and TB07.

Special Instructions: 2°C temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #
Analysis Key: GC\_Alcohol=07-Alcohols, Anions=07-Anions, Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, OilGrease=07-Oil & Grease, HEM, WetChem=07-Phosphorous\_NO3-NO2-Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen, TDS=07-Solids, Total Dissolved, TSS=07-Solids, Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains one row of data.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

Page 2 of 2

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SECT 301.101 (General) - 40 CFR 136.1 (NEW) CHAIN OF CUSTODY RECORD No: 3-012812-103308-0008
Date Shipped: 1/28/2012 Site #: A3TA Lab: EPA R3 Laboratory
Carrier Name: FedEx Case Complete: False Lab Contact:
Airbill No: 797998070399 Lab Phone: 410.305.3032

Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains two rows of sample data.

Special Instructions: 2°C Temp Blank
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #
Analysis Key: GC\_Alcohol=07-Alcohols, Anions=07-Anions\_Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, Oil/Grease=07-Oil & Grease, HEM, WetChem=07-Phosphorous\_NO3-, NO2-, Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen, TDS=07-Solids, Total Dissolved, TSS=07-Solids, Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains one row of chain of custody data.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center
Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater

Project #: DAS R33907

Page 1 of 2

SELECTION OF CHAIN OF CUSTODY RECORD

No: 3-012812-110206-0010

Date Shipped: 1/28/2012

Site #: A37A

Lab: EPA R3 Laboratory

Carrier Name: FedEx

Case Complete: False

Lab Contact:

Airbill No: 7979 9807 0436

Lab Phone: 410.305.3032

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Table with 8 columns: Sample #, Matrix/Sampler, Coll. Method, Analysis/Turnaround, Tag/Preservative/Bottles, Station Location, Collected, For Lab Use Only. Contains sample data for HW14-P, HW14-PF, and TB04.

Special Instructions: No Len P Blank (cooler was packed with ice)
Shipment for Case Complete? N
Samples Transferred From Chain of Custody #

Analysis Key: GC\_Alcohol=07-Alcohols, Anions=07-Anions, Method 300.0, Glycols=07-Glycols+2-Butoxyethanol+EGME, Metals=07-Metals, WetChem=07-Phosphorous, NO3-, NO2-, Tot-N, SVOC+TIC=07-Semivolatiles (TCL+TICs+EGME+1-Methylnaphthalen, TDS=07-Solids, Total Dissolved, TSS=07-Solids, Total Suspended, VOC+TIC=07-Volatiles+Acrylonitrile (TCL+TICs), F-Metals=07-Metals Dissolved

Table with 12 columns: Items/Reason, Relinquished by, Date, Received by, Date, Time, Items/Reason, Relinquished By, Date, Received by, Date, Time. Contains a single entry for item 19.



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center  
Office of Analytical Services and Quality Assurance  
701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB01	<b>Lab ID:</b> 1201013-01
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/23/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/26/12	01/27/12 15:22	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/26/12	01/27/12 15:22	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/26/12	01/27/12 15:22	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/26/12	01/27/12 15:22	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/26/12	01/27/12 15:22	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Acenaphthylene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Acetophenone	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Anthracene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Atrazine	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Benzaldehyde	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Benzo(a)anthracene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Benzo(a)pyrene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
1,1-Biphenyl	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	0.358	B, J	5.00	1	01/25/12	01/26/12 16:46	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Carbazole	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Caprolactam	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
4-Chloroaniline	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
2-Chloronaphthalene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
2-Chlorophenol	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Chrysene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
Dibenzofuran	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB01 Lab ID: 1201013-01
Sample Matrix: Water Date Collected: 01/23/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various chemical compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc., with their respective results and flags.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB01	<b>Lab ID:</b> 1201013-01
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/23/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/25/12	01/26/12 16:46	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	69.7		70 %	21-110	01/25/12	01/26/12 16:46	R3QA201
<i>Surrogate: Phenol-d5</i>	82.8		83 %	10-110	01/25/12	01/26/12 16:46	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	37.5		75 %	35-114	01/25/12	01/26/12 16:46	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	36.6		73 %	43-116	01/25/12	01/26/12 16:46	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	82.7		83 %	10-123	01/25/12	01/26/12 16:46	R3QA201
<i>Surrogate: Terphenyl-d14</i>	44.2		88 %	33-141	01/25/12	01/26/12 16:46	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.8	J	2.0	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Benzene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Bromoform	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
<b>Chloroform</b>	7.5		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/27/12	01/27/12 14:48	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB01 Lab ID: 1201013-01
Sample Matrix: Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB01 Lab ID: 1201013-01
Sample Matrix: Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW19	<b>Lab ID:</b> 1201013-03
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/23/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/26/12	01/27/12 15:36	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/26/12	01/27/12 15:36	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/26/12	01/27/12 15:36	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/26/12	01/27/12 15:36	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/26/12	01/27/12 15:36	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Acenaphthylene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Acetophenone	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Anthracene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Atrazine	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Benzaldehyde	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Benzo(a)anthracene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Benzo(a)pyrene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
1,1-Biphenyl	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	0.214	B, J	5.00	1	01/26/12	01/27/12 16:59	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
<b>Butyl benzyl phthalate</b>	0.040	B, J	5.00	1	01/26/12	01/27/12 16:59	R3QA201
Carbazole	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Caprolactam	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
4-Chloroaniline	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
2-Chloronaphthalene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
2-Chlorophenol	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Chrysene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
Dibenzofuran	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/26/12	01/27/12 16:59	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19 Lab ID: 1201013-03
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Lists various organic compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19 Lab ID: 1201013-03
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Row 1: 2,4,6-Trichlorophenol, U, 5.00, 1, 01/26/12, 01/27/12 16:59, R3QA201

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 2-Fluorophenol, Phenol-d5, Nitrobenzene-d5, 2-Fluorobiphenyl, 2,4,6-Tribromophenol, Terphenyl-d14

Volatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Lists various organic compounds like Acetone, Benzene, Bromobenzene, etc.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19 Lab ID: 1201013-03
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19 Lab ID: 1201013-03
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater
Station ID: HW19-P
Sample Matrix: Drinking Water
Project #: DAS R33907
Lab ID: 1201013-05
Date Collected: 01/23/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19-P Lab ID: 1201013-05
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, Fluoranthene, and Pyrene.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW19-P	<b>Lab ID:</b> 1201013-05
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/23/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/26/12	01/27/12 17:49	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	56.8		57 %	21-110	01/26/12	01/27/12 17:49	R3QA201
<i>Surrogate: Phenol-d5</i>	66.8		67 %	10-110	01/26/12	01/27/12 17:49	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	29.3		59 %	35-114	01/26/12	01/27/12 17:49	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	30.3		61 %	43-116	01/26/12	01/27/12 17:49	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	69.9		70 %	10-123	01/26/12	01/27/12 17:49	R3QA201
<i>Surrogate: Terphenyl-d14</i>	36.1		72 %	33-141	01/26/12	01/27/12 17:49	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.3	B, J	2.0	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Benzene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Bromoform	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Chloroform	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/27/12	01/27/12 16:37	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19-P Lab ID: 1201013-05
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW19-P Lab ID: 1201013-05
Sample Matrix: Drinking Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB02	<b>Lab ID:</b> 1201013-07
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/24/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/30/12	01/30/12 09:00	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/30/12	01/30/12 09:00	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/30/12	01/30/12 09:00	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/30/12	01/30/12 09:00	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/30/12	01/30/12 09:00	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Acenaphthylene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Acetophenone	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Anthracene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Atrazine	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Benzaldehyde	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Benzo(a)anthracene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Benzo(a)pyrene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
1,1-Biphenyl	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	0.451	B, J	5.00	1	01/29/12	01/30/12 22:10	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
<b>Butyl benzyl phthalate</b>	0.079	B, J	5.00	1	01/29/12	01/30/12 22:10	R3QA201
Carbazole	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Caprolactam	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
4-Chloroaniline	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
2-Chloronaphthalene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
2-Chlorophenol	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Chrysene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
Dibenzofuran	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB02 Lab ID: 1201013-07
Sample Matrix: Water Date Collected: 01/24/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, Di-n-butyl phthalate, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB02	<b>Lab ID:</b> 1201013-07
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/24/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/30/12 22:10	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	%Recovery	Limits			
Surrogate: 2-Fluorophenol	85.1		85 %	21-110	01/29/12	01/30/12 22:10	R3QA201
Surrogate: Phenol-d5	97.0		97 %	10-110	01/29/12	01/30/12 22:10	R3QA201
Surrogate: Nitrobenzene-d5	43.5		87 %	35-114	01/29/12	01/30/12 22:10	R3QA201
Surrogate: 2-Fluorobiphenyl	43.0		86 %	43-116	01/29/12	01/30/12 22:10	R3QA201
Surrogate: 2,4,6-Tribromophenol	96.9		97 %	10-123	01/29/12	01/30/12 22:10	R3QA201
Surrogate: Terphenyl-d14	49.9		100 %	33-141	01/29/12	01/30/12 22:10	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.9	J	2.0	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Benzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>Bromodichloromethane</b>	0.06	J	0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Bromoform	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>Chloroform</b>	7.6		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB02	<b>Lab ID:</b> 1201013-07
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/24/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags Qualifiers	Quantitation		Dilution	Prepared	Analyzed	Method/SOP#
	ug/L		Limit					
1,2-Dibromo-3-chloropropane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Dibromomethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2-Dichloroethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Ethylbenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Freon 113	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
2-Hexanone	U		2.0		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Isopropylbenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Methyl Acetate	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Methylcyclohexane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.0		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
n-Propylbenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Styrene	U		1.0		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Tetrachloroethene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>Toluene</b>	0.6		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5		1	01/27/12	01/27/12 15:16	CLP trace/R3QA210



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Office of Analytical Services and Quality Assurance  
701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB02	<b>Lab ID:</b> 1201013-07
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/24/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
<b>Trichloroethene</b>	1.5		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.2	J	1.0	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<b>o-Xylene</b>	0.07	J	1.0	1	01/27/12	01/27/12 15:16	CLP trace/R3QA210

**Surrogates**

Analyte	Result ug/L	Flags Qualifiers	%Recovery	%Recovery Limits	Prepared	Analyzed	Method/SOP#
<i>Surrogate: 4-Bromofluorobenzene</i>	3.920		<b>98 %</b>	86-115	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.240		<b>106 %</b>	76-114	01/27/12	01/27/12 15:16	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.930		<b>98 %</b>	88-110	01/27/12	01/27/12 15:16	CLP trace/R3QA210



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW04	<b>Lab ID:</b> 1201013-09
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/24/2012

**Alcohols  
Targets**

Analyte	Result ug/mL	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
1-Butanol	U		10.0	1	01/30/12	01/30/12 09:14	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/30/12	01/30/12 09:14	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/30/12	01/30/12 09:14	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/30/12	01/30/12 09:14	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/30/12	01/30/12 09:14	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
Acenaphthene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Acenaphthylene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Acetophenone	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Anthracene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Atrazine	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Benzaldehyde	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Benzo(a)anthracene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
<b>Benzo(a)pyrene</b>	0.049	J	5.00	1	01/29/12	01/30/12 23:00	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
1,1-Biphenyl	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	0.437	B, J	5.00	1	01/29/12	01/30/12 23:00	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
<b>Butyl benzyl phthalate</b>	0.069	B, J	5.00	1	01/29/12	01/30/12 23:00	R3QA201
Carbazole	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Caprolactam	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
4-Chloroaniline	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
2-Chloronaphthalene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
2-Chlorophenol	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Chrysene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
Dibenzofuran	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW04 Lab ID: 1201013-09
Sample Matrix: Drinking Water Date Collected: 01/24/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW04	<b>Lab ID:</b> 1201013-09
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/24/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/30/12 23:00	R3QA201

**Surrogates**

Analyte	Result ug/mL	Flags Qualifiers	%Recovery %Recovery	%Recovery Limits	Prepared	Analyzed	Method/SOP#
Surrogate: 2-Fluorophenol	91.3		91 %	21-110	01/29/12	01/30/12 23:00	R3QA201
Surrogate: Phenol-d5	94.9		95 %	10-110	01/29/12	01/30/12 23:00	R3QA201
Surrogate: Nitrobenzene-d5	42.3		85 %	35-114	01/29/12	01/30/12 23:00	R3QA201
Surrogate: 2-Fluorobiphenyl	42.9		86 %	43-116	01/29/12	01/30/12 23:00	R3QA201
Surrogate: 2,4,6-Tribromophenol	98.4		98 %	10-123	01/29/12	01/30/12 23:00	R3QA201
Surrogate: Terphenyl-d14	48.4		97 %	33-141	01/29/12	01/30/12 23:00	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
Acetone	0.3	B, J	2.0	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Benzene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Bromoform	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Chloroform	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/27/12	01/27/12 17:04	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW04 Lab ID: 1201013-09
Sample Matrix: Drinking Water Date Collected: 01/24/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW04 Lab ID: 1201013-09
Sample Matrix: Drinking Water Date Collected: 01/24/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: TB01 Lab ID: 1201013-11
Sample Matrix: Water Date Collected: 01/23/2012

Volatile Organic Compounds
Targets

Table with 9 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acetone, Benzene, Bromobenzene, Bromochloromethane, Bromodichloromethane, Bromoform, Bromomethane, 2-Butanone, sec-Butylbenzene, tert-Butylbenzene, n-Butylbenzene, Carbon disulfide, Carbon Tetrachloride, Chlorobenzene, Chlorodibromomethane, Chloroethane, Chloroform, Chloromethane, 2-Chlorotoluene, 4-Chlorotoluene, Cyclohexane, 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane (EDB), Dibromomethane, 1,2-Dichlorobenzene, 1,3-Dichlorobenzene, 1,4-Dichlorobenzene, Dichlorodifluoromethane, 1,1-Dichloroethane, 1,2-Dichloroethane, 1,1-Dichloroethene, cis-1,2-Dichloroethene, trans-1,2-Dichloroethene, 1,2-Dichloropropane, 1,3-Dichloropropane, 2,2-Dichloropropane, 1,1-Dichloropropene, cis-1,3-Dichloropropene, trans-1,3-Dichloropropene, Ethylbenzene.



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB01	<b>Lab ID:</b> 1201013-11
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/23/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.0		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Styrene	U		1.0	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<b>Toluene</b>	0.6		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<b>Trichloroethene</b>	2.7		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.2	J	1.0	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<b>o-Xylene</b>	0.07	J	1.0	1	01/27/12	01/27/12 15:43	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.810		<b>95 %</b>	86-115	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.260		<b>106 %</b>	76-114	01/27/12	01/27/12 15:43	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.930		<b>98 %</b>	88-110	01/27/12	01/27/12 15:43	CLP trace/R3QA210



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB03	<b>Lab ID:</b> 1201013-12
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/25/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/30/12	01/30/12 09:27	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/30/12	01/30/12 09:27	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/30/12	01/30/12 09:27	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/30/12	01/30/12 09:27	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/30/12	01/30/12 09:27	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Acenaphthylene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Acetophenone	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Anthracene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Atrazine	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Benzaldehyde	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Benzo(a)anthracene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Benzo(a)pyrene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
1,1-Biphenyl	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	0.369	B, J	5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Carbazole	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Caprolactam	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Chloroaniline	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2-Chloronaphthalene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2-Chlorophenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Chrysene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Dibenzofuran	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

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 701 Mapes Road  
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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB03	<b>Lab ID:</b> 1201013-12
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/25/2012

Semivolatile Organic Compounds  
 Targets (Continued)

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Diethyl phthalate</b>	0.043	B, J	5.00	1	01/29/12	01/30/12 23:51	R3QA201
2,4-Dichlorophenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2,4-Dimethylphenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Dimethyl phthalate	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2,4-Dinitrophenol	U	UJ	5.00	1	01/29/12	01/30/12 23:51	R3QA201
<b>Di-n-butyl phthalate</b>	0.397	B, J	5.00	1	01/29/12	01/30/12 23:51	R3QA201
4,6-Dinitro-2-methylphenol	U	UJ	10.0	1	01/29/12	01/30/12 23:51	R3QA201
2,4-Dinitrotoluene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2,6-Dinitrotoluene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Di-n-octyl phthalate	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Fluoranthene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Fluorene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Hexachlorobenzene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Hexachlorobutadiene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Hexachlorocyclopentadiene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Hexachloroethane	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Indeno(1,2,3-cd)pyrene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Isophorone	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2-Methoxyethanol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
<b>1-Methylnaphthalene</b>	0.024	J	5.00	1	01/29/12	01/30/12 23:51	R3QA201
<b>2-Methylnaphthalene</b>	0.027	J	5.00	1	01/29/12	01/30/12 23:51	R3QA201
2-Methylphenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Methylphenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
<b>Naphthalene</b>	0.332	J	5.00	1	01/29/12	01/30/12 23:51	R3QA201
2-Nitroaniline	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
3-Nitroaniline	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Nitroaniline	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Nitrobenzene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2-Nitrophenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
4-Nitrophenol	U		10.0	1	01/29/12	01/30/12 23:51	R3QA201
N-Nitrosodimethylamine	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
N-Nitroso-di-n-propylamine	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
N-Nitrosodiphenylamine	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Pentachlorophenol	U	UJ	5.00	1	01/29/12	01/30/12 23:51	R3QA201
Phenanthrene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Phenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
Pyrene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
1,2,4,5-Tetrachlorobenzene	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2,3,4,6-Tetrachlorophenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201
2,4,5-Trichlorophenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB03	<b>Lab ID:</b> 1201013-12
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/25/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/30/12 23:51	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	%Recovery	Limits			
Surrogate: 2-Fluorophenol	87.4		87 %	21-110	01/29/12	01/30/12 23:51	R3QA201
Surrogate: Phenol-d5	91.3		91 %	10-110	01/29/12	01/30/12 23:51	R3QA201
Surrogate: Nitrobenzene-d5	40.5		81 %	35-114	01/29/12	01/30/12 23:51	R3QA201
Surrogate: 2-Fluorobiphenyl	40.5		81 %	43-116	01/29/12	01/30/12 23:51	R3QA201
Surrogate: 2,4,6-Tribromophenol	91.5		92 %	10-123	01/29/12	01/30/12 23:51	R3QA201
Surrogate: Terphenyl-d14	46.8		94 %	33-141	01/29/12	01/30/12 23:51	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	1.0	J	2.0	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Benzene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Bromoform	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
<b>Chloroform</b>	8.3		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/30/12	01/30/12 15:20	CLP trace/R3QA210



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB03 Lab ID: 1201013-12
Sample Matrix: Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include various chemical compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB03 Lab ID: 1201013-12
Sample Matrix: Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater
Station ID: HW02
Sample Matrix: Drinking Water
Project #: DAS R33907
Lab ID: 1201013-13
Date Collected: 01/25/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine, Diethyl phthalate.



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW02	<b>Lab ID:</b> 1201013-13
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

Semivolatile Organic Compounds  
Targets (Continued)

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4-Dichlorophenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2,4-Dimethylphenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Dimethyl phthalate	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2,4-Dinitrophenol	U	UJ	5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>Di-n-butyl phthalate</b>	0.709	B, J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
4,6-Dinitro-2-methylphenol	U	UJ	10.0	1	01/29/12	01/31/12 00:41	R3QA201
2,4-Dinitrotoluene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2,6-Dinitrotoluene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>Di-n-octyl phthalate</b>	0.193	J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>Fluoranthene</b>	0.198	J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>Fluorene</b>	0.077	J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>Hexachlorobenzene</b>	0.200	J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
Hexachlorobutadiene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Hexachlorocyclopentadiene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Hexachloroethane	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Indeno(1,2,3-cd)pyrene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Isophorone	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2-Methoxyethanol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
1-Methylnaphthalene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2-Methylnaphthalene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2-Methylphenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
4-Methylphenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Naphthalene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2-Nitroaniline	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
3-Nitroaniline	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
4-Nitroaniline	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Nitrobenzene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2-Nitrophenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
4-Nitrophenol	U		10.0	1	01/29/12	01/31/12 00:41	R3QA201
N-Nitrosodimethylamine	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
N-Nitroso-di-n-propylamine	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>N-Nitrosodiphenylamine</b>	0.034	J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
Pentachlorophenol	U	UJ	5.00	1	01/29/12	01/31/12 00:41	R3QA201
Phenanthrene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
Phenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
<b>Pyrene</b>	0.183	J	5.00	1	01/29/12	01/31/12 00:41	R3QA201
1,2,4,5-Tetrachlorobenzene	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2,3,4,6-Tetrachlorophenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2,4,5-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 00:41	R3QA201



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701 Mapes Road  
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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW02	<b>Lab ID:</b> 1201013-13
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

Semivolatile Organic Compounds

Surrogates

Analyte	Result ug/mL	Flags Qualifiers	%Recovery	%Recovery Limits	Prepared	Analyzed	Method/SOP#
Surrogate: 2-Fluorophenol	76.2		76 %	21-110	01/29/12	01/31/12 00:41	R3QA201
Surrogate: Phenol-d5	86.5		87 %	10-110	01/29/12	01/31/12 00:41	R3QA201
Surrogate: Nitrobenzene-d5	39.5		79 %	35-114	01/29/12	01/31/12 00:41	R3QA201
Surrogate: 2-Fluorobiphenyl	39.7		79 %	43-116	01/29/12	01/31/12 00:41	R3QA201
Surrogate: 2,4,6-Tribromophenol	84.9		85 %	10-123	01/29/12	01/31/12 00:41	R3QA201
Surrogate: Terphenyl-d14	44.1		88 %	33-141	01/29/12	01/31/12 00:41	R3QA201

Volatile Organic Compounds

Targets

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
Acetone	0.3	B, J	2.0	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Benzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Bromoform	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Chloroform	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2-Dibromo-3-chloropropane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW02	<b>Lab ID:</b> 1201013-13
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Dichlorodifluoromethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2-Dichloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Freon 113	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Methylene Chloride	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Naphthalene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Styrene	U		1.0	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Toluene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Trichloroethene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/30/12	01/30/12 16:43	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW02 Lab ID: 1201013-13
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW02z	<b>Lab ID:</b> 1201013-14
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/30/12	01/30/12 09:55	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/30/12	01/30/12 09:55	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/30/12	01/30/12 09:55	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/30/12	01/30/12 09:55	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/30/12	01/30/12 09:55	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Acenaphthylene</b>	0.013	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Acetophenone	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Anthracene</b>	0.231	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Atrazine	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Benzaldehyde	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Benzo(a)anthracene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Benzo(a)pyrene</b>	0.196	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Benzo(b)fluoranthene</b>	0.150	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Benzo(ghi)perylene</b>	0.211	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Benzo(k)fluoranthene</b>	0.317	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
1,1-Biphenyl	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	2.68	B, J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>4-Bromophenyl phenyl ether</b>	0.179	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Butyl benzyl phthalate</b>	0.351	B, J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Carbazole</b>	0.287	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Caprolactam	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
4-Chloroaniline	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2-Chloronaphthalene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2-Chlorophenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>4-Chlorophenyl phenyl ether</b>	0.096	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Chrysene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Dibenzofuran</b>	0.038	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Diethyl phthalate</b>	0.257	B, J	5.00	1	01/29/12	01/31/12 01:32	R3QA201



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW02z	<b>Lab ID:</b> 1201013-14
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

Semivolatile Organic Compounds  
Targets (Continued)

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4-Dichlorophenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2,4-Dimethylphenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Dimethyl phthalate</b>	0.149	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
2,4-Dinitrophenol	U	UJ	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Di-n-butyl phthalate</b>	0.848	B, J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
4,6-Dinitro-2-methylphenol	U	UJ	10.0	1	01/29/12	01/31/12 01:32	R3QA201
<b>2,4-Dinitrotoluene</b>	0.130	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
2,6-Dinitrotoluene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Di-n-octyl phthalate</b>	0.281	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Fluoranthene</b>	0.268	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Fluorene</b>	0.098	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Hexachlorobenzene</b>	0.217	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Hexachlorobutadiene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Hexachlorocyclopentadiene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Hexachloroethane	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Indeno(1,2,3-cd)pyrene</b>	0.205	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Isophorone	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2-Methoxyethanol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
1-Methylnaphthalene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2-Methylnaphthalene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2-Methylphenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
4-Methylphenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
Naphthalene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>2-Nitroaniline</b>	0.074	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>3-Nitroaniline</b>	0.119	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>4-Nitroaniline</b>	0.166	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Nitrobenzene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2-Nitrophenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>4-Nitrophenol</b>	0.136	J	10.0	1	01/29/12	01/31/12 01:32	R3QA201
N-Nitrosodimethylamine	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
N-Nitroso-di-n-propylamine	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>N-Nitrosodiphenylamine</b>	0.171	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Pentachlorophenol	U	UJ	5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Phenanthrene</b>	0.234	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
Phenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
<b>Pyrene</b>	0.257	J	5.00	1	01/29/12	01/31/12 01:32	R3QA201
1,2,4,5-Tetrachlorobenzene	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2,3,4,6-Tetrachlorophenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2,4,5-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 01:32	R3QA201



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW02z	<b>Lab ID:</b> 1201013-14
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

Semivolatile Organic Compounds

Surrogates

Analyte	Result ug/mL	Flags Qualifiers	%Recovery	%Recovery Limits	Prepared	Analyzed	Method/SOP#
Surrogate: 2-Fluorophenol	76.4		76 %	21-110	01/29/12	01/31/12 01:32	R3QA201
Surrogate: Phenol-d5	88.8		89 %	10-110	01/29/12	01/31/12 01:32	R3QA201
Surrogate: Nitrobenzene-d5	39.5		79 %	35-114	01/29/12	01/31/12 01:32	R3QA201
Surrogate: 2-Fluorobiphenyl	40.1		80 %	43-116	01/29/12	01/31/12 01:32	R3QA201
Surrogate: 2,4,6-Tribromophenol	91.1		91 %	10-123	01/29/12	01/31/12 01:32	R3QA201
Surrogate: Terphenyl-d14	45.1		90 %	33-141	01/29/12	01/31/12 01:32	R3QA201

Volatile Organic Compounds

Targets

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
Acetone	U		2.0	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Benzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Bromoform	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Chloroform	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
1,2-Dibromo-3-chloropropane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/30/12	01/30/12 17:10	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW02z Lab ID: 1201013-14
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like Dichlorodifluoromethane, 1,1-Dichloroethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW02z Lab ID: 1201013-14
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW01 Lab ID: 1201013-15
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW01 Lab ID: 1201013-15
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW01	<b>Lab ID:</b> 1201013-15
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 02:22	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	%Recovery	Limits			
Surrogate: 2-Fluorophenol	80.1		80 %	21-110	01/29/12	01/31/12 02:22	R3QA201
Surrogate: Phenol-d5	91.3		91 %	10-110	01/29/12	01/31/12 02:22	R3QA201
Surrogate: Nitrobenzene-d5	40.5		81 %	35-114	01/29/12	01/31/12 02:22	R3QA201
Surrogate: 2-Fluorobiphenyl	40.1		80 %	43-116	01/29/12	01/31/12 02:22	R3QA201
Surrogate: 2,4,6-Tribromophenol	93.5		94 %	10-123	01/29/12	01/31/12 02:22	R3QA201
Surrogate: Terphenyl-d14	46.0		92 %	33-141	01/29/12	01/31/12 02:22	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acetone	1.1	B, J	2.0	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Benzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Bromoform	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Chloroform	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW01 Lab ID: 1201013-15
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW01	<b>Lab ID:</b> 1201013-15
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Volatile Organic Compounds  
 Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Trichloroethene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
m-Xylene/p-Xylene	U		1.0	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210
o-Xylene	U		1.0	1	01/30/12	01/30/12 17:38	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.760		<b>94 %</b>	<i>86-115</i>	01/30/12	01/30/12 17:38	<i>CLP trace/R3QA210</i>
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.080		<b>102 %</b>	<i>76-114</i>	01/30/12	01/30/12 17:38	<i>CLP trace/R3QA210</i>
<i>Surrogate: Toluene-d8</i>	3.640		<b>91 %</b>	<i>88-110</i>	01/30/12	01/30/12 17:38	<i>CLP trace/R3QA210</i>



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW05 Lab ID: 1201013-16
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, and 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW05 Lab ID: 1201013-16
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW05	<b>Lab ID:</b> 1201013-16
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 03:12	R3QA201

**Surrogates**

Analyte	Result ug/mL	Flags Qualifiers	%Recovery %Recovery	%Recovery Limits	Prepared	Analyzed	Method/SOP#
<i>Surrogate: 2-Fluorophenol</i>	73.6		<b>74 %</b>	<i>21-110</i>	01/29/12	01/31/12 03:12	<i>R3QA201</i>
<i>Surrogate: Phenol-d5</i>	89.1		<b>89 %</b>	<i>10-110</i>	01/29/12	01/31/12 03:12	<i>R3QA201</i>
<i>Surrogate: Nitrobenzene-d5</i>	40.0		<b>80 %</b>	<i>35-114</i>	01/29/12	01/31/12 03:12	<i>R3QA201</i>
<i>Surrogate: 2-Fluorobiphenyl</i>	40.7		<b>81 %</b>	<i>43-116</i>	01/29/12	01/31/12 03:12	<i>R3QA201</i>
<i>Surrogate: 2,4,6-Tribromophenol</i>	90.0		<b>90 %</b>	<i>10-123</i>	01/29/12	01/31/12 03:12	<i>R3QA201</i>
<i>Surrogate: Terphenyl-d14</i>	45.5		<b>91 %</b>	<i>33-141</i>	01/29/12	01/31/12 03:12	<i>R3QA201</i>

**Volatile Organic Compounds  
Targets**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
<b>Acetone</b>	0.4	B, J	2.0	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Benzene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Bromoform	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Chloroform	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/30/12	01/30/12 18:05	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW05 Lab ID: 1201013-16
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW05 Lab ID: 1201013-16
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater
Station ID: HW06
Sample Matrix: Drinking Water
Project #: DAS R33907
Lab ID: 1201013-17
Date Collected: 01/26/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW06 Lab ID: 1201013-17
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, Fluoranthene, Pyrene, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW06	<b>Lab ID:</b> 1201013-17
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 04:02	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	81.0		<b>81 %</b>	<i>21-110</i>	01/29/12	01/31/12 04:02	<i>R3QA201</i>
<i>Surrogate: Phenol-d5</i>	92.6		<b>93 %</b>	<i>10-110</i>	01/29/12	01/31/12 04:02	<i>R3QA201</i>
<i>Surrogate: Nitrobenzene-d5</i>	41.4		<b>83 %</b>	<i>35-114</i>	01/29/12	01/31/12 04:02	<i>R3QA201</i>
<i>Surrogate: 2-Fluorobiphenyl</i>	41.4		<b>83 %</b>	<i>43-116</i>	01/29/12	01/31/12 04:02	<i>R3QA201</i>
<i>Surrogate: 2,4,6-Tribromophenol</i>	93.3		<b>93 %</b>	<i>10-123</i>	01/29/12	01/31/12 04:02	<i>R3QA201</i>
<i>Surrogate: Terphenyl-d14</i>	35.8		<b>72 %</b>	<i>33-141</i>	01/29/12	01/31/12 04:02	<i>R3QA201</i>

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.3	B, J	2.0	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Benzene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Bromoform	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Chloroform	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/30/12	01/30/12 18:32	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW06 Lab ID: 1201013-17
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW06 Lab ID: 1201013-17
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: TB02 Lab ID: 1201013-25
Sample Matrix: Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets

Table with 9 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acetone, Benzene, Bromobenzene, Bromochloromethane, Bromodichloromethane, Bromoform, Bromomethane, 2-Butanone, sec-Butylbenzene, tert-Butylbenzene, n-Butylbenzene, Carbon disulfide, Carbon Tetrachloride, Chlorobenzene, Chlorodibromomethane, Chloroethane, Chloroform, Chloromethane, 2-Chlorotoluene, 4-Chlorotoluene, Cyclohexane, 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane (EDB), Dibromomethane, 1,2-Dichlorobenzene, 1,3-Dichlorobenzene, 1,4-Dichlorobenzene, Dichlorodifluoromethane, 1,1-Dichloroethane, 1,2-Dichloroethane, 1,1-Dichloroethene, cis-1,2-Dichloroethene, trans-1,2-Dichloroethene, 1,2-Dichloropropane, 1,3-Dichloropropane, 2,2-Dichloropropane, 1,1-Dichloropropene, cis-1,3-Dichloropropene, trans-1,3-Dichloropropene, Ethylbenzene.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB02	<b>Lab ID:</b> 1201013-25
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/25/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.1		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Styrene	U		1.0	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<b>Toluene</b>	0.6		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<b>Trichloroethene</b>	2.1		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.2	J	1.0	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<b>o-Xylene</b>	0.06	J	1.0	1	01/30/12	01/30/12 15:48	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.910		<b>98 %</b>	86-115	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.220		<b>106 %</b>	76-114	01/30/12	01/30/12 15:48	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.750		<b>94 %</b>	88-110	01/30/12	01/30/12 15:48	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: TB03 Lab ID: 1201013-26
Sample Matrix: Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets

Table with 9 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acetone, Benzene, Bromobenzene, Bromochloromethane, Bromodichloromethane, Bromoform, Bromomethane, 2-Butanone, sec-Butylbenzene, tert-Butylbenzene, n-Butylbenzene, Carbon disulfide, Carbon Tetrachloride, Chlorobenzene, Chlorodibromomethane, Chloroethane, Chloroform, Chloromethane, 2-Chlorotoluene, 4-Chlorotoluene, Cyclohexane, 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane (EDB), Dibromomethane, 1,2-Dichlorobenzene, 1,3-Dichlorobenzene, 1,4-Dichlorobenzene, Dichlorodifluoromethane, 1,1-Dichloroethane, 1,2-Dichloroethane, 1,1-Dichloroethene, cis-1,2-Dichloroethene, trans-1,2-Dichloroethene, 1,2-Dichloropropane, 1,3-Dichloropropane, 2,2-Dichloropropane, 1,1-Dichloropropene, cis-1,3-Dichloropropene, trans-1,3-Dichloropropene, Ethylbenzene.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB03	<b>Lab ID:</b> 1201013-26
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/25/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.1		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Styrene	U		1.0	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<b>Toluene</b>	0.6		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<b>Trichloroethene</b>	2.4		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.2	J	1.0	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<b>o-Xylene</b>	0.07	J	1.0	1	01/30/12	01/30/12 16:15	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.780		<b>94 %</b>	86-115	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.140		<b>104 %</b>	76-114	01/30/12	01/30/12 16:15	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.710		<b>93 %</b>	88-110	01/30/12	01/30/12 16:15	CLP trace/R3QA210



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center  
Office of Analytical Services and Quality Assurance  
701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW08a	<b>Lab ID:</b> 1201013-28
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/30/12	01/30/12 11:17	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/30/12	01/30/12 11:17	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/30/12	01/30/12 11:17	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/30/12	01/30/12 11:17	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/30/12	01/30/12 11:17	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Acenaphthylene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Acetophenone	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
<b>Anthracene</b>	0.056	J	5.00	1	01/29/12	01/31/12 04:53	R3QA201
Atrazine	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Benzaldehyde	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Benzo(a)anthracene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Benzo(a)pyrene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
1,1-Biphenyl	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	1.24	B, J	5.00	1	01/29/12	01/31/12 04:53	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
<b>Butyl benzyl phthalate</b>	0.105	B, J	5.00	1	01/29/12	01/31/12 04:53	R3QA201
Carbazole	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Caprolactam	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
4-Chloroaniline	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
2-Chloronaphthalene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
2-Chlorophenol	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
<b>4-Chlorophenyl phenyl ether</b>	0.029	J	5.00	1	01/29/12	01/31/12 04:53	R3QA201
Chrysene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
Dibenzofuran	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW08a Lab ID: 1201013-28
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc., with their respective results and limits.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW08a	<b>Lab ID:</b> 1201013-28
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/29/12	01/31/12 04:53	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	72.0		72 %	21-110	01/29/12	01/31/12 04:53	R3QA201
<i>Surrogate: Phenol-d5</i>	87.4		87 %	10-110	01/29/12	01/31/12 04:53	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	39.2		78 %	35-114	01/29/12	01/31/12 04:53	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	39.5		79 %	43-116	01/29/12	01/31/12 04:53	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	92.8		93 %	10-123	01/29/12	01/31/12 04:53	R3QA201
<i>Surrogate: Terphenyl-d14</i>	45.0		90 %	33-141	01/29/12	01/31/12 04:53	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.8	B, J	2.0	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW08a	<b>Lab ID:</b> 1201013-28
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/25/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
1,2-Dibromo-3-chloropropane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,2-Dichloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Freon 113	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Methylene Chloride	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Naphthalene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
Toluene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 14:59	CLP trace/R3QA210



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW08a Lab ID: 1201013-28
Sample Matrix: Drinking Water Date Collected: 01/25/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB04	<b>Lab ID:</b> 1201013-29
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/26/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/31/12	01/31/12 18:32	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/31/12	01/31/12 18:32	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/31/12	01/31/12 18:32	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/31/12	01/31/12 18:32	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/31/12	01/31/12 18:32	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Acenaphthylene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
<b>Acetophenone</b>	0.043	J	5.00	1	01/31/12	02/01/12 19:31	R3QA201
Anthracene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Atrazine	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Benzaldehyde	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Benzo(a)anthracene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Benzo(a)pyrene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
1,1-Biphenyl	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	2.02	B, J	5.00	1	01/31/12	02/01/12 19:31	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Carbazole	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Caprolactam	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
4-Chloroaniline	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
2-Chloronaphthalene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
2-Chlorophenol	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Chrysene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
Dibenzofuran	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201



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Office of Analytical Services and Quality Assurance
701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB04 Lab ID: 1201013-29
Sample Matrix: Water Date Collected: 01/26/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, Di-n-butyl phthalate, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB04	<b>Lab ID:</b> 1201013-29
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/26/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 19:31	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	75.2		75 %	21-110	01/31/12	02/01/12 19:31	R3QA201
<i>Surrogate: Phenol-d5</i>	81.5		82 %	10-110	01/31/12	02/01/12 19:31	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	34.5		69 %	35-114	01/31/12	02/01/12 19:31	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	34.2		68 %	43-116	01/31/12	02/01/12 19:31	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	82.9		83 %	10-123	01/31/12	02/01/12 19:31	R3QA201
<i>Surrogate: Terphenyl-d14</i>	41.9		84 %	33-141	01/31/12	02/01/12 19:31	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	1.3	J	2.0	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
<b>Bromodichloromethane</b>	0.2	J	0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
<b>Chloroform</b>	3.0		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB04	<b>Lab ID:</b> 1201013-29
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/26/2012

Volatile Organic Compounds  
Targets (Continued)

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
1,2-Dibromo-3-chloropropane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
<b>1,2-Dichloroethane</b>	0.1	J	0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Freon 113	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
<b>Methylene Chloride</b>	2.0		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
<b>Naphthalene</b>	0.2	J	0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
<b>Toluene</b>	0.5	J	0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 15:29	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB04 Lab ID: 1201013-29
Sample Matrix: Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater
Station ID: FB05
Sample Matrix: Water
Project #: DAS R33907
Lab ID: 1201013-30
Date Collected: 01/27/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB05	<b>Lab ID:</b> 1201013-30
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/27/2012

Semivolatile Organic Compounds  
Targets (Continued)

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Diethyl phthalate</b>	0.344	B, J	5.00	1	01/31/12	02/01/12 20:22	R3QA201
2,4-Dichlorophenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2,4-Dimethylphenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Dimethyl phthalate	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2,4-Dinitrophenol	U	UJ	5.00	1	01/31/12	02/01/12 20:22	R3QA201
<b>Di-n-butyl phthalate</b>	0.455	B, J	5.00	1	01/31/12	02/01/12 20:22	R3QA201
4,6-Dinitro-2-methylphenol	U	UJ	10.0	1	01/31/12	02/01/12 20:22	R3QA201
2,4-Dinitrotoluene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2,6-Dinitrotoluene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Di-n-octyl phthalate	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Fluoranthene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Fluorene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Hexachlorobenzene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Hexachlorobutadiene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Hexachlorocyclopentadiene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Hexachloroethane	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Indeno(1,2,3-cd)pyrene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Isophorone	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2-Methoxyethanol	U	UJ	5.00	1	01/31/12	02/01/12 20:22	R3QA201
<b>1-Methylnaphthalene</b>	0.021	J	5.00	1	01/31/12	02/01/12 20:22	R3QA201
<b>2-Methylnaphthalene</b>	0.027	J	5.00	1	01/31/12	02/01/12 20:22	R3QA201
2-Methylphenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
4-Methylphenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
<b>Naphthalene</b>	0.291	J	5.00	1	01/31/12	02/01/12 20:22	R3QA201
2-Nitroaniline	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
3-Nitroaniline	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
4-Nitroaniline	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Nitrobenzene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2-Nitrophenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
4-Nitrophenol	U		10.0	1	01/31/12	02/01/12 20:22	R3QA201
N-Nitrosodimethylamine	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
N-Nitroso-di-n-propylamine	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
N-Nitrosodiphenylamine	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Pentachlorophenol	U	UJ	5.00	1	01/31/12	02/01/12 20:22	R3QA201
Phenanthrene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Phenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
Pyrene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
1,2,4,5-Tetrachlorobenzene	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2,3,4,6-Tetrachlorophenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201
2,4,5-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB05	<b>Lab ID:</b> 1201013-30
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/27/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 20:22	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	86.2		86 %	21-110	01/31/12	02/01/12 20:22	R3QA201
<i>Surrogate: Phenol-d5</i>	94.7		95 %	10-110	01/31/12	02/01/12 20:22	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	40.0		80 %	35-114	01/31/12	02/01/12 20:22	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	39.1		78 %	43-116	01/31/12	02/01/12 20:22	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	92.2		92 %	10-123	01/31/12	02/01/12 20:22	R3QA201
<i>Surrogate: Terphenyl-d14</i>	45.0		90 %	33-141	01/31/12	02/01/12 20:22	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	1.6	J	2.0	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
<b>Bromodichloromethane</b>	0.2	J	0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
<b>Chloroform</b>	2.8		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> FB05	<b>Lab ID:</b> 1201013-30
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/27/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
1,2-Dibromo-3-chloropropane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
<b>1,2-Dichloroethane</b>	0.07	J	0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Freon 113	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
<b>Methylene Chloride</b>	2.0		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
<b>Naphthalene</b>	0.2	J	0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
<b>Toluene</b>	0.9		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 15:59	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: FB05 Lab ID: 1201013-30
Sample Matrix: Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW24	<b>Lab ID:</b> 1201013-31
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/31/12	01/31/12 18:59	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/31/12	01/31/12 18:59	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/31/12	01/31/12 18:59	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/31/12	01/31/12 18:59	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/31/12	01/31/12 18:59	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Acenaphthylene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Acetophenone	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Anthracene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Atrazine	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Benzaldehyde	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Benzo(a)anthracene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Benzo(a)pyrene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
1,1-Biphenyl	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	2.48	B, J	5.00	1	01/31/12	02/01/12 21:13	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Carbazole	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Caprolactam	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
4-Chloroaniline	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
2-Chloronaphthalene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
2-Chlorophenol	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Chrysene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
Dibenzofuran	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW24 Lab ID: 1201013-31
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 9 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW24	<b>Lab ID:</b> 1201013-31
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 21:13	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	101		<b>101 %</b>	21-110	01/31/12	02/01/12 21:13	R3QA201
<i>Surrogate: Phenol-d5</i>	110		<b>110 %</b>	10-110	01/31/12	02/01/12 21:13	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	46.8		<b>94 %</b>	35-114	01/31/12	02/01/12 21:13	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	47.6		<b>95 %</b>	43-116	01/31/12	02/01/12 21:13	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	114		<b>114 %</b>	10-123	01/31/12	02/01/12 21:13	R3QA201
<i>Surrogate: Terphenyl-d14</i>	50.0		<b>100 %</b>	33-141	01/31/12	02/01/12 21:13	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.5	B, J	2.0	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
<b>Carbon disulfide</b>	0.1	J	0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 18:27	CLP trace/R3QA210



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701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW24 Lab ID: 1201013-31
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW24 Lab ID: 1201013-31
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW24-P	<b>Lab ID:</b> 1201013-32
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/31/12	01/31/12 19:41	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/31/12	01/31/12 19:41	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/31/12	01/31/12 19:41	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/31/12	01/31/12 19:41	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/31/12	01/31/12 19:41	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Acenaphthylene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Acetophenone	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Anthracene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Atrazine	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Benzaldehyde	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Benzo(a)anthracene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Benzo(a)pyrene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
1,1-Biphenyl	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	1.76	B, J	5.00	1	01/31/12	02/01/12 22:04	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
<b>Butyl benzyl phthalate</b>	0.057	B, J	5.00	1	01/31/12	02/01/12 22:04	R3QA201
Carbazole	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Caprolactam	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
4-Chloroaniline	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
2-Chloronaphthalene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
2-Chlorophenol	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Chrysene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
Dibenzofuran	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW24-P Lab ID: 1201013-32
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW24-P	<b>Lab ID:</b> 1201013-32
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 22:04	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	78.1		<b>78 %</b>	<i>21-110</i>	01/31/12	02/01/12 22:04	<i>R3QA201</i>
<i>Surrogate: Phenol-d5</i>	84.9		<b>85 %</b>	<i>10-110</i>	01/31/12	02/01/12 22:04	<i>R3QA201</i>
<i>Surrogate: Nitrobenzene-d5</i>	36.1		<b>72 %</b>	<i>35-114</i>	01/31/12	02/01/12 22:04	<i>R3QA201</i>
<i>Surrogate: 2-Fluorobiphenyl</i>	37.5		<b>75 %</b>	<i>43-116</i>	01/31/12	02/01/12 22:04	<i>R3QA201</i>
<i>Surrogate: 2,4,6-Tribromophenol</i>	87.1		<b>87 %</b>	<i>10-123</i>	01/31/12	02/01/12 22:04	<i>R3QA201</i>
<i>Surrogate: Terphenyl-d14</i>	45.0		<b>90 %</b>	<i>33-141</i>	01/31/12	02/01/12 22:04	<i>R3QA201</i>

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.7	B, J	2.0	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
<b>Carbon disulfide</b>	0.05	J	0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 18:56	CLP trace/R3QA210



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW24-P Lab ID: 1201013-32
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and analysis dates.



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW24-P Lab ID: 1201013-32
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, and o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, and Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW12	<b>Lab ID:</b> 1201013-33
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/31/12	01/31/12 19:54	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/31/12	01/31/12 19:54	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/31/12	01/31/12 19:54	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/31/12	01/31/12 19:54	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/31/12	01/31/12 19:54	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Acenaphthylene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Acetophenone	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Anthracene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Atrazine	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Benzaldehyde	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Benzo(a)anthracene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Benzo(a)pyrene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
1,1-Biphenyl	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	1.32	B, J	5.00	1	01/31/12	02/01/12 22:55	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Carbazole	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Caprolactam	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
4-Chloroaniline	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
2-Chloronaphthalene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
2-Chlorophenol	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Chrysene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
Dibenzofuran	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW12 Lab ID: 1201013-33
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Lists various organic compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc., with their respective results and limits.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW12	<b>Lab ID:</b> 1201013-33
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 22:55	R3QA201

**Surrogates**

Analyte	Result ug/L	Flags Qualifiers	%Recovery %Recovery	Limit	Prepared	Analyzed	Method/SOP#
<i>Surrogate: 2-Fluorophenol</i>	44.2		<b>44 %</b>	21-110	01/31/12	02/01/12 22:55	R3QA201
<i>Surrogate: Phenol-d5</i>	50.0		<b>50 %</b>	10-110	01/31/12	02/01/12 22:55	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	24.8		<b>50 %</b>	35-114	01/31/12	02/01/12 22:55	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	27.1		<b>54 %</b>	43-116	01/31/12	02/01/12 22:55	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	69.7		<b>70 %</b>	10-123	01/31/12	02/01/12 22:55	R3QA201
<i>Surrogate: Terphenyl-d14</i>	37.8		<b>76 %</b>	33-141	01/31/12	02/01/12 22:55	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
<b>Acetone</b>	1.8	B, J	2.0	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
<b>Chloroethane</b>	0.2	J	0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
<b>Chloromethane</b>	0.6		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

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Office of Analytical Services and Quality Assurance  
701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW12	<b>Lab ID:</b> 1201013-33
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
1,2-Dibromo-3-chloropropane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,2-Dichloroethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Freon 113	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Methylene Chloride	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Naphthalene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
Toluene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 19:24	CLP trace/R3QA210



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW12 Lab ID: 1201013-33
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW17	<b>Lab ID:</b> 1201013-34
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/31/12	01/31/12 20:08	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/31/12	01/31/12 20:08	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/31/12	01/31/12 20:08	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/31/12	01/31/12 20:08	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/31/12	01/31/12 20:08	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Acenaphthylene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Acetophenone	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Anthracene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Atrazine	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Benzaldehyde	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Benzo(a)anthracene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Benzo(a)pyrene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
1,1-Biphenyl	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	1.16	B, J	5.00	1	01/31/12	02/01/12 23:46	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Carbazole	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Caprolactam	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
4-Chloroaniline	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
2-Chloronaphthalene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
2-Chlorophenol	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Chrysene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
Dibenzofuran	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201



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 Office of Analytical Services and Quality Assurance  
 701 Mapes Road  
 Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW17	<b>Lab ID:</b> 1201013-34
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

Semivolatile Organic Compounds  
 Targets (Continued)

Analyte	Result	Flags	Quantitation		Dilution	Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit					
<b>Diethyl phthalate</b>	0.273	B, J	5.00		1	01/31/12	02/01/12 23:46	R3QA201
2,4-Dichlorophenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2,4-Dimethylphenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Dimethyl phthalate	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2,4-Dinitrophenol	U	UJ	5.00		1	01/31/12	02/01/12 23:46	R3QA201
<b>Di-n-butyl phthalate</b>	0.814	B, J	5.00		1	01/31/12	02/01/12 23:46	R3QA201
4,6-Dinitro-2-methylphenol	U	UJ	10.0		1	01/31/12	02/01/12 23:46	R3QA201
2,4-Dinitrotoluene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2,6-Dinitrotoluene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Di-n-octyl phthalate	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Fluoranthene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Fluorene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Hexachlorobenzene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Hexachlorobutadiene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Hexachlorocyclopentadiene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Hexachloroethane	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Indeno(1,2,3-cd)pyrene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Isophorone	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2-Methoxyethanol	U	UJ	5.00		1	01/31/12	02/01/12 23:46	R3QA201
1-Methylnaphthalene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2-Methylnaphthalene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2-Methylphenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
4-Methylphenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Naphthalene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2-Nitroaniline	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
3-Nitroaniline	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
4-Nitroaniline	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Nitrobenzene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2-Nitrophenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
4-Nitrophenol	U		10.0		1	01/31/12	02/01/12 23:46	R3QA201
N-Nitrosodimethylamine	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
N-Nitroso-di-n-propylamine	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
N-Nitrosodiphenylamine	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Pentachlorophenol	U	UJ	5.00		1	01/31/12	02/01/12 23:46	R3QA201
Phenanthrene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Phenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
Pyrene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
1,2,4,5-Tetrachlorobenzene	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2,3,4,6-Tetrachlorophenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201
2,4,5-Trichlorophenol	U		5.00		1	01/31/12	02/01/12 23:46	R3QA201



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW17	<b>Lab ID:</b> 1201013-34
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/27/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/01/12 23:46	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	46.3		46 %	21-110	01/31/12	02/01/12 23:46	R3QA201
<i>Surrogate: Phenol-d5</i>	56.9		57 %	10-110	01/31/12	02/01/12 23:46	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	27.6		55 %	35-114	01/31/12	02/01/12 23:46	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	29.3		59 %	43-116	01/31/12	02/01/12 23:46	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	68.4		68 %	10-123	01/31/12	02/01/12 23:46	R3QA201
<i>Surrogate: Terphenyl-d14</i>	35.8		72 %	33-141	01/31/12	02/01/12 23:46	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	1.2	B, J	2.0	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 19:52	CLP trace/R3QA210



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW17 Lab ID: 1201013-34
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW17 Lab ID: 1201013-34
Sample Matrix: Drinking Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW14	<b>Lab ID:</b> 1201013-35
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Alcohols  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/mL	Qualifiers	Limit	Dilution			
1-Butanol	U		10.0	1	01/31/12	01/31/12 20:22	EPA 8015D/R3QA203
2-Butanol	U		10.0	1	01/31/12	01/31/12 20:22	EPA 8015D/R3QA203
Ethanol	U		10.0	1	01/31/12	01/31/12 20:22	EPA 8015D/R3QA203
Methanol	U		10.0	1	01/31/12	01/31/12 20:22	EPA 8015D/R3QA203
1-Propanol	U		10.0	1	01/31/12	01/31/12 20:22	EPA 8015D/R3QA203

**Semivolatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Acenaphthene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Acenaphthylene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Acetophenone	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Anthracene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Atrazine	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Benzaldehyde	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Benzo(a)anthracene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Benzo(a)pyrene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Benzo(b)fluoranthene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Benzo(ghi)perylene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Benzo(k)fluoranthene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
1,1-Biphenyl	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Bis(2-chloroethoxy)methane	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Bis(2-chloroethyl)ether	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Bis(2-chloroisopropyl)ether	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
<b>Bis(2-ethylhexyl)phthalate</b>	1.45	B, J	5.00	1	01/31/12	02/02/12 00:37	R3QA201
4-Bromophenyl phenyl ether	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Butyl benzyl phthalate	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Carbazole	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Caprolactam	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
4-Chloroaniline	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
4-Chloro-3-methylphenol	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
2-Chloronaphthalene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
2-Chlorophenol	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
4-Chlorophenyl phenyl ether	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Chrysene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Dibenz(a,h)anthracene	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
Dibenzofuran	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201
3,3'-Dichlorobenzidine	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW14 Lab ID: 1201013-35
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 9 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW14	<b>Lab ID:</b> 1201013-35
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/02/12 00:37	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
Surrogate: 2-Fluorophenol	61.7		62 %	21-110	01/31/12	02/02/12 00:37	R3QA201
Surrogate: Phenol-d5	69.6		70 %	10-110	01/31/12	02/02/12 00:37	R3QA201
Surrogate: Nitrobenzene-d5	34.9		70 %	35-114	01/31/12	02/02/12 00:37	R3QA201
Surrogate: 2-Fluorobiphenyl	35.7		71 %	43-116	01/31/12	02/02/12 00:37	R3QA201
Surrogate: 2,4,6-Tribromophenol	73.3		73 %	10-123	01/31/12	02/02/12 00:37	R3QA201
Surrogate: Terphenyl-d14	41.7		83 %	33-141	01/31/12	02/02/12 00:37	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	0.6	B, J	2.0	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
<b>Bromoform</b>	0.2	J	0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
2-Butanone	U		2.0	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 20:20	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW14 Lab ID: 1201013-35
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW14 Lab ID: 1201013-35
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Trichloroethene, Trichlorofluoromethane, 1,2,3-Trichloropropane, 1,2,4-Trimethylbenzene, 1,3,5-Trimethylbenzene, Vinyl acetate, Vinyl chloride, m-Xylene/p-Xylene, o-Xylene.

Surrogates

Table with 8 columns: Analyte, Result, Flags, %Recovery, %Recovery Limits, Prepared, Analyzed, Method/SOP#. Rows include Surrogate: 4-Bromofluorobenzene, Surrogate: 1,2-Dichloroethane-d4, Surrogate: Toluene-d8.



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW14-P Lab ID: 1201013-36
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Alcohols
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Semivolatile Organic Compounds
Targets

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acenaphthene, Acenaphthylene, Acetophenone, Anthracene, Atrazine, Benzaldehyde, Benzo(a)anthracene, Benzo(a)pyrene, Benzo(b)fluoranthene, Benzo(ghi)perylene, Benzo(k)fluoranthene, 1,1-Biphenyl, Bis(2-chloroethoxy)methane, Bis(2-chloroethyl)ether, Bis(2-chloroisopropyl)ether, Bis(2-ethylhexyl)phthalate, 4-Bromophenyl phenyl ether, Butyl benzyl phthalate, Carbazole, Caprolactam, 4-Chloroaniline, 4-Chloro-3-methylphenol, 2-Chloronaphthalene, 2-Chlorophenol, 4-Chlorophenyl phenyl ether, Chrysene, Dibenz(a,h)anthracene, Dibenzofuran, 3,3'-Dichlorobenzidine.



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701 Mapes Road
Fort Meade, Maryland 20755-5350



Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW14-P Lab ID: 1201013-36
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Semivolatile Organic Compounds
Targets (Continued)

Table with 9 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include various compounds like Diethyl phthalate, 2,4-Dichlorophenol, etc.



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW14-P	<b>Lab ID:</b> 1201013-36
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Semivolatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
2,4,6-Trichlorophenol	U		5.00	1	01/31/12	02/02/12 01:27	R3QA201

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 2-Fluorophenol</i>	76.7		77 %	21-110	01/31/12	02/02/12 01:27	R3QA201
<i>Surrogate: Phenol-d5</i>	84.6		85 %	10-110	01/31/12	02/02/12 01:27	R3QA201
<i>Surrogate: Nitrobenzene-d5</i>	35.4		71 %	35-114	01/31/12	02/02/12 01:27	R3QA201
<i>Surrogate: 2-Fluorobiphenyl</i>	35.9		72 %	43-116	01/31/12	02/02/12 01:27	R3QA201
<i>Surrogate: 2,4,6-Tribromophenol</i>	79.7		80 %	10-123	01/31/12	02/02/12 01:27	R3QA201
<i>Surrogate: Terphenyl-d14</i>	41.0		82 %	33-141	01/31/12	02/02/12 01:27	R3QA201

**Volatile Organic Compounds  
Targets**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
<b>Acetone</b>	81.1	K	2.0	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
<b>2-Butanone</b>	1.0	B, J	2.0	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Chloroform	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: HW14-P Lab ID: 1201013-36
Sample Matrix: Drinking Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets (Continued)

Table with 8 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows list various compounds like 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane, etc., with their respective results and limits.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> HW14-P	<b>Lab ID:</b> 1201013-36
<b>Sample Matrix:</b> Drinking Water	<b>Date Collected:</b> 01/26/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Trichloroethene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
m-Xylene/p-Xylene	U		1.0	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210
o-Xylene	U		1.0	1	01/31/12	01/31/12 20:48	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.880		<b>97 %</b>	<i>86-115</i>	01/31/12	01/31/12 20:48	<i>CLP trace/R3QA210</i>
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.250		<b>106 %</b>	<i>76-114</i>	01/31/12	01/31/12 20:48	<i>CLP trace/R3QA210</i>
<i>Surrogate: Toluene-d8</i>	3.570		<b>89 %</b>	<i>88-110</i>	01/31/12	01/31/12 20:48	<i>CLP trace/R3QA210</i>



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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: TB05 Lab ID: 1201013-45
Sample Matrix: Water Date Collected: 01/26/2012

Volatile Organic Compounds
Targets

Table with 9 columns: Analyte, Result (ug/L), Flags/Qualifiers, Quantitation (Limit), Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acetone, Benzene, Bromobenzene, Bromochloromethane, Bromodichloromethane, Bromoform, Bromomethane, 2-Butanone, sec-Butylbenzene, tert-Butylbenzene, n-Butylbenzene, Carbon disulfide, Carbon Tetrachloride, Chlorobenzene, Chlorodibromomethane, Chloroethane, Chloroform, Chloromethane, 2-Chlorotoluene, 4-Chlorotoluene, Cyclohexane, 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane (EDB), Dibromomethane, 1,2-Dichlorobenzene, 1,3-Dichlorobenzene, 1,4-Dichlorobenzene, Dichlorodifluoromethane, 1,1-Dichloroethane, 1,2-Dichloroethane, 1,1-Dichloroethene, cis-1,2-Dichloroethene, trans-1,2-Dichloroethene, 1,2-Dichloropropane, 1,3-Dichloropropane, 2,2-Dichloropropane, 1,1-Dichloropropene, cis-1,3-Dichloropropene, trans-1,3-Dichloropropene, Ethylbenzene.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB05	<b>Lab ID:</b> 1201013-45
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/26/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.1		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<b>Toluene</b>	0.6		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<b>Trichloroethene</b>	2.0		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.1	J	1.0	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<b>o-Xylene</b>	0.06	J	1.0	1	01/31/12	01/31/12 16:59	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.870		<b>97 %</b>	86-115	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.360		<b>109 %</b>	76-114	01/31/12	01/31/12 16:59	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.760		<b>94 %</b>	88-110	01/31/12	01/31/12 16:59	CLP trace/R3QA210



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701 Mapes Road  
Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB07	<b>Lab ID:</b> 1201013-46
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/27/2012

**Volatile Organic Compounds  
Targets**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
<b>Acetone</b>	1.3	J	2.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Bromodichloromethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>2-Butanone</b>	0.4	J	2.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>Chloroform</b>	7.7		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2-Dibromo-3-chloropropane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2-Dichloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210



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Fort Meade, Maryland 20755-5350



<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB07	<b>Lab ID:</b> 1201013-46
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/27/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.1		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>Toluene</b>	0.6		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>Trichloroethene</b>	1.5		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.1	J	1.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210
o-Xylene	U		1.0	1	01/31/12	01/31/12 16:29	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.780		<b>94 %</b>	86-115	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.510		<b>113 %</b>	76-114	01/31/12	01/31/12 16:29	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.680		<b>92 %</b>	88-110	01/31/12	01/31/12 16:29	CLP trace/R3QA210



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701 Mapes Road
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Site Name: Dimock Residential Groundwater Project #: DAS R33907
Station ID: TB06 Lab ID: 1201013-47
Sample Matrix: Water Date Collected: 01/27/2012

Volatile Organic Compounds
Targets

Table with 9 columns: Analyte, Result, Flags, Quantitation, Dilution, Prepared, Analyzed, Method/SOP#. Rows include Acetone, Benzene, Bromobenzene, Bromochloromethane, Bromodichloromethane, Bromoform, Bromomethane, 2-Butanone, sec-Butylbenzene, tert-Butylbenzene, n-Butylbenzene, Carbon disulfide, Carbon Tetrachloride, Chlorobenzene, Chlorodibromomethane, Chloroethane, Chloroform, Chloromethane, 2-Chlorotoluene, 4-Chlorotoluene, Cyclohexane, 1,2-Dibromo-3-chloropropane, 1,2-Dibromoethane (EDB), Dibromomethane, 1,2-Dichlorobenzene, 1,3-Dichlorobenzene, 1,4-Dichlorobenzene, Dichlorodifluoromethane, 1,1-Dichloroethane, 1,2-Dichloroethane, 1,1-Dichloroethene, cis-1,2-Dichloroethene, trans-1,2-Dichloroethene, 1,2-Dichloropropane, 1,3-Dichloropropane, 2,2-Dichloropropane, 1,1-Dichloropropene, cis-1,3-Dichloropropene, trans-1,3-Dichloropropene, Ethylbenzene.



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB06	<b>Lab ID:</b> 1201013-47
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/27/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.1		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<b>Toluene</b>	0.5		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<b>Trichloroethene</b>	1.6		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.1	J	1.0	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210
o-Xylene	U		1.0	1	01/31/12	01/31/12 17:58	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.890		<b>97 %</b>	86-115	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.590	A	<b>115 %</b>	76-114	01/31/12	01/31/12 17:58	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.620		<b>90 %</b>	88-110	01/31/12	01/31/12 17:58	CLP trace/R3QA210



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701 Mapes Road  
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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB04	<b>Lab ID:</b> 1201013-48
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/26/2012

**Volatile Organic Compounds  
Targets**

Analyte	Result ug/L	Flags Qualifiers	Quantitation Limit	Dilution	Prepared	Analyzed	Method/SOP#
<b>Acetone</b>	1.4	J	2.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Benzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Bromobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Bromochloromethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>Bromodichloromethane</b>	0.05	J	0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Bromoform	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Bromomethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>2-Butanone</b>	0.5	J	2.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
sec-Butylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
tert-Butylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
n-Butylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Carbon disulfide	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Carbon Tetrachloride	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Chlorobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Chlorodibromomethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Chloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>Chloroform</b>	8.2		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Chloromethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
2-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
4-Chlorotoluene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Cyclohexane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2-Dibromo-3-chloropropane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2-Dibromoethane (EDB)	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Dibromomethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,3-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,4-Dichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Dichlorodifluoromethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1-Dichloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2-Dichloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1-Dichloroethene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
cis-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
trans-1,2-Dichloroethene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,3-Dichloropropane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
2,2-Dichloropropane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1-Dichloropropene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
cis-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
trans-1,3-Dichloropropene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Ethylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210



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<b>Site Name:</b> Dimock Residential Groundwater	<b>Project #:</b> DAS R33907
<b>Station ID:</b> TB04	<b>Lab ID:</b> 1201013-48
<b>Sample Matrix:</b> Water	<b>Date Collected:</b> 01/26/2012

**Volatile Organic Compounds  
Targets (Continued)**

Analyte	Result	Flags	Quantitation		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	Limit	Dilution			
Freon 113	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Hexachlorobutadiene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
2-Hexanone	U		2.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Isopropylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
p-Isopropyltoluene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Methyl Acetate	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Methylcyclohexane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Methyl-tert-butyl ether	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>Methylene Chloride</b>	1.2		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
4-Methyl-2-pentanone	U		2.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>Naphthalene</b>	0.3	J	0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
n-Propylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Styrene	U		1.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1,2,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1,1,2-Tetrachloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Tetrachloroethene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>Toluene</b>	0.7		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2,3-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2,4-Trichlorobenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1,1-Trichloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,1,2-Trichloroethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>Trichloroethene</b>	2.2		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Trichlorofluoromethane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2,3-Trichloropropane	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,2,4-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
1,3,5-Trimethylbenzene	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Vinyl acetate	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
Vinyl chloride	U		0.5	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>m-Xylene/p-Xylene</b>	0.1	J	1.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<b>o-Xylene</b>	0.08	J	1.0	1	01/31/12	01/31/12 17:29	CLP trace/R3QA210

**Surrogates**

Analyte	Result	Flags	%Recovery		Prepared	Analyzed	Method/SOP#
	ug/L	Qualifiers	%Recovery	Limits			
<i>Surrogate: 4-Bromofluorobenzene</i>	3.760		<b>94 %</b>	86-115	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<i>Surrogate: 1,2-Dichloroethane-d4</i>	4.380		<b>110 %</b>	76-114	01/31/12	01/31/12 17:29	CLP trace/R3QA210
<i>Surrogate: Toluene-d8</i>	3.640		<b>91 %</b>	88-110	01/31/12	01/31/12 17:29	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-01					
<b>Station ID:</b>	FB01					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/23/2012					
74367-33-2	Propanoic acid, 2-methyl-, 2,2-dimethyl-1-(2-h	6.66	T	6.43	01/26/12 16:46	R3QA201
82304-66-3	7,9-Di-tert-butyl-1-oxaspiro(4,5)deca-6,9-dien	3.18	T	9.32	01/26/12 16:46	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-01					
<b>Station ID:</b>	FB01					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/23/2012					
	None Detected	0.0			01/27/12 14:48	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

Tentatively Identified Compound (TIC) Report  
Semivolatile Organic Compounds

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-03					
<b>Station ID:</b>	HW19					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/23/2012					
	None Detected	0.00			01/27/12 16:59	R3QA201

Volatile Organic Compounds

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-03					
<b>Station ID:</b>	HW19					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/23/2012					
	None Detected	0.0			01/27/12 16:10	CLP trace/R3QA210

Semivolatile Organic Compounds

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-05					
<b>Station ID:</b>	HW19-P					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/23/2012					
	None Detected	0.00			01/27/12 17:49	R3QA201

Volatile Organic Compounds

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-05					
<b>Station ID:</b>	HW19-P					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/23/2012					
	None Detected	0.0			01/27/12 16:37	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-07					
<b>Station ID:</b>	FB02					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/24/2012					
NA	unknown	7.77	T	6.43	01/30/12 22:10	R3QA201
82304-66-3	7,9-Di-tert-butyl-1-oxaspiro(4,5)deca-6,9-dien	3.50	T	9.32	01/30/12 22:10	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-07					
<b>Station ID:</b>	FB02					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/24/2012					
	None Detected	0.0			01/27/12 15:16	CLP trace/R3QA210



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

Tentatively Identified Compound (TIC) Report
Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes sample details (Lab ID, Station ID, Sample Matrix, Collected) and four rows of compound data.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes sample details and one row of compound data showing 'None Detected'.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes sample details and one row of compound data showing 'None Detected'.



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Project #: DAS R33907

**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-12					
<b>Station ID:</b>	FB03					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/25/2012					
NA	unknown	8.54	T	6.43	01/30/12 23:51	R3QA201
82304-66-3	7,9-Di-tert-butyl-1-oxaspiro(4,5)deca-6,9-dien	5.46	T	9.32	01/30/12 23:51	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-12					
<b>Station ID:</b>	FB03					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/25/2012					
	None Detected	0.0			01/30/12 15:20	CLP trace/R3QA210



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Tentatively Identified Compound (TIC) Report
Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected date.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected date.

Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected date.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected date.



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Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info. Data row: 7517-36-4 Phthalic acid, monocyclohexyl ester 3.21 T 12.74 01/31/12 02:22 R3QA201

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info. Data row: 74-98-6 Propane 0.4 T 1.17 01/30/12 17:38 CLP trace/R3QA210

Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info. Data row: None Detected 0.00 01/31/12 03:12 R3QA201

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info. Data row: None Detected 0.0 01/30/12 18:05 CLP trace/R3QA210



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Tentatively Identified Compound (TIC) Report
Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.



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Tentatively Identified Compound (TIC) Report
Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-26, Station ID: TB03, Sample Matrix: Water, Collected: 01/25/2012, and a row for 'None Detected' with result 0.0.

Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-28, Station ID: HW08a, Sample Matrix: Drinking Water, Collected: 01/25/2012, and rows for 4-Acetamidoacetophenone, 2(1H)-Quinolinone, 3-methyl-, and two unknown compounds.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-28, Station ID: HW08a, Sample Matrix: Drinking Water, Collected: 01/25/2012, and a row for Propane with result 0.8.



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**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-29					
<b>Station ID:</b>	FB04					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/26/2012					
NA	unknown (01)	11.8	T	6.43	02/01/12 19:31	R3QA201
82304-66-3	7,9-Di-tert-butyl-1-oxaspiro(4,5)deca-6,9-dien	3.16	T	9.32	02/01/12 19:31	R3QA201
NA	unknown (02)	3.75	T	17.02	02/01/12 19:31	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-29					
<b>Station ID:</b>	FB04					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/26/2012					
	None Detected	0.0			01/31/12 15:29	CLP trace/R3QA210



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**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-30					
<b>Station ID:</b>	FB05					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/27/2012					
NA	unknown (01)	17.0	T	6.43	02/01/12 20:22	R3QA201
82304-66-3	7,9-Di-tert-butyl-1-oxaspiro(4,5)deca-6,9-dien	3.28	T	9.32	02/01/12 20:22	R3QA201
NA	unknown (02)	3.49	T	17.02	02/01/12 20:22	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-30					
<b>Station ID:</b>	FB05					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/27/2012					
	None Detected	0.0			01/31/12 15:59	CLP trace/R3QA210



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**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-31					
<b>Station ID:</b>	HW24					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/27/2012					
	None Detected	0.00			02/01/12 21:13	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-31					
<b>Station ID:</b>	HW24					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/27/2012					
420-56-4	Trimethylsilyl fluoride	1.9	T	1.60	01/31/12 18:27	CLP trace/R3QA210



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Tentatively Identified Compound (TIC) Report

Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected date. Lists compounds like Octacosane, Nonacosane, and Hentriacontane.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected date. Lists Trimethylsilyl fluoride.



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**Tentatively Identified Compound (TIC) Report**  
**Semivolatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-33					
<b>Station ID:</b>	HW12					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/26/2012					
	None Detected	0.00			02/01/12 22:55	R3QA201

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-33					
<b>Station ID:</b>	HW12					
<b>Sample Matrix:</b>	Drinking Water					
<b>Collected:</b>	01/26/2012					
74-98-6	Propane	30.9	T	1.17	01/31/12 19:24	CLP trace/R3QA210
75-28-5	Isobutane	2.6	T	1.33	01/31/12 19:24	CLP trace/R3QA210
106-97-8	Butane	4.1	T	1.48	01/31/12 19:24	CLP trace/R3QA210



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Tentatively Identified Compound (TIC) Report
Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-34, Station ID: HW17, Sample Matrix: Drinking Water, Collected: 01/27/2012, and data row: None Detected, 0.00, 02/01/12 23:46, R3QA201.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-34, Station ID: HW17, Sample Matrix: Drinking Water, Collected: 01/27/2012, and data row: None Detected, 0.0, 01/31/12 19:52, CLP trace/R3QA210.

Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-35, Station ID: HW14, Sample Matrix: Drinking Water, Collected: 01/26/2012, and data row: None Detected, 0.00, 02/02/12 00:37, R3QA201.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID: 1201013-35, Station ID: HW14, Sample Matrix: Drinking Water, Collected: 01/26/2012, and data row: None Detected, 0.0, 01/31/12 20:20, CLP trace/R3QA210.



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Tentatively Identified Compound (TIC) Report
Semivolatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.

Volatile Organic Compounds

Table with 7 columns: CAS Number, Compound, Result (ug/L), Analyte Qualifiers, Retention Time, Analyzed, Method/SOP#. Includes Lab ID, Station ID, Sample Matrix, and Collected info.



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**Tentatively Identified Compound (TIC) Report**  
**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-47					
<b>Station ID:</b>	TB06					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/27/2012					
	None Detected	0.0			01/31/12 17:58	CLP trace/R3QA210

**Volatile Organic Compounds**

CAS Number	Compound	Result ug/L	Analyte Qualifiers	Retention Time	Analyzed	Method/SOP#
<b>Lab ID:</b>	1201013-48					
<b>Station ID:</b>	TB04					
<b>Sample Matrix:</b>	Water					
<b>Collected:</b>	01/26/2012					
	None Detected	0.0			01/31/12 17:29	CLP trace/R3QA210



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QC Data
Alcohols

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22701 - Alcohols

Blank (BA22701-BLK1)

Prepared: 01/26/12 08:00 Analyzed: 01/27/12 14:54

Table with 4 columns: Analyte, Result, Quantitation Limit, Units. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

LCS (BA22701-BS1)

Prepared: 01/26/12 08:00 Analyzed: 01/27/12 15:08

Table with 6 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Matrix Spike (BA22701-MS1)

Source: 1201013-03

Prepared: 01/26/12 08:00 Analyzed: 01/27/12 15:49

Table with 6 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Matrix Spike Dup (BA22701-MSD1)

Source: 1201013-03

Prepared: 01/26/12 08:00 Analyzed: 01/27/12 16:03

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.



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QC Data
Alcohols

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA23001 - Alcohols

Blank (BA23001-BLK1)

Prepared: 01/30/12 07:02 Analyzed: 01/30/12 08:32

Table with 4 columns: Analyte, Result, Quantitation Limit, Units. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

LCS (BA23001-BS1)

Prepared: 01/30/12 07:02 Analyzed: 01/30/12 08:46

Table with 7 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, %REC, %REC Limits. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Matrix Spike (BA23001-MS1)

Source: 1201013-17

Prepared: 01/30/12 07:02 Analyzed: 01/30/12 10:50

Table with 8 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.

Matrix Spike Dup (BA23001-MSD1)

Source: 1201013-17

Prepared: 01/30/12 07:02 Analyzed: 01/30/12 11:03

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit. Rows for 1-Butanol, 2-Butanol, Ethanol, Methanol, 1-Propanol.



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**QC Data**  
**Alcohols**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BA23103 - Alcohols**

**Blank (BA23103-BLK1)**

Prepared: 01/31/12 09:16 Analyzed: 01/31/12 18:05

1-Butanol	U	10.0	ug/mL							
2-Butanol	U	10.0	"							
Ethanol	U	10.0	"							
Methanol	U	10.0	"							
1-Propanol	U	10.0	"							

**LCS (BA23103-BS1)**

Prepared: 01/31/12 09:16 Analyzed: 01/31/12 18:18

1-Butanol	100	10.0	ug/mL	100.00		100	70-130			
2-Butanol	96.3	10.0	"	100.00		96	70-130			
Ethanol	97.5	10.0	"	100.00		97	70-130			
Methanol	91.5	10.0	"	100.00		91	70-130			
1-Propanol	97.5	10.0	"	100.00		98	70-130			

**Matrix Spike (BA23103-MS1)**

Source: 1201013-31

Prepared: 01/31/12 09:16 Analyzed: 01/31/12 19:13

1-Butanol	105	10.0	ug/mL	100.00	0.00	105	70-130			
2-Butanol	101	10.0	"	100.00	0.00	101	70-130			
Ethanol	101	10.0	"	100.00	0.00	101	70-130			
Methanol	94.1	10.0	"	100.00	0.00	94	70-130			
1-Propanol	102	10.0	"	100.00	0.00	102	70-130			

**Matrix Spike Dup (BA23103-MSD1)**

Source: 1201013-31

Prepared: 01/31/12 09:16 Analyzed: 01/31/12 19:27

1-Butanol	104	10.0	ug/mL	100.00	0.00	104	70-130	2	25	
2-Butanol	99.8	10.0	"	100.00	0.00	100	70-130	1	25	
Ethanol	96.4	10.0	"	100.00	0.00	96	70-130	4	25	
Methanol	90.8	10.0	"	100.00	0.00	91	70-130	4	25	
1-Propanol	99.3	10.0	"	100.00	0.00	99	70-130	3	25	



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QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22504 - EPA 3520C SVOC

Blank (BA22504-BLK1)

Prepared: 01/25/12 12:43 Analyzed: 01/26/12 14:14

Main data table listing various chemical compounds (e.g., Acenaphthene, Benzaldehyde, etc.) with their corresponding results (U, 0.408, 0.088, 0.966) and units (ug/L).



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**QC Data**  
**Semivolatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BA22504 - EPA 3520C SVOC**

**Blank (BA22504-BLK1)**

Prepared: 01/25/12 12:43 Analyzed: 01/26/12 14:14

Hexachlorobenzene	U	5.00	ug/L							
Hexachlorobutadiene	U	5.00	"							
Hexachlorocyclopentadiene	U	5.00	"							
Hexachloroethane	U	5.00	"							
Indeno(1,2,3-cd)pyrene	U	5.00	"							
Isophorone	U	5.00	"							
2-Methoxyethanol	U	5.00	"							
1-Methylnaphthalene	U	5.00	"							
2-Methylnaphthalene	U	5.00	"							
2-Methylphenol	U	5.00	"							
4-Methylphenol	U	5.00	"							
Naphthalene	U	5.00	"							
2-Nitroaniline	U	5.00	"							
3-Nitroaniline	U	5.00	"							
4-Nitroaniline	U	5.00	"							
Nitrobenzene	U	5.00	"							
2-Nitrophenol	U	5.00	"							
4-Nitrophenol	U	10.0	"							
N-Nitrosodimethylamine	U	5.00	"							
N-Nitroso-di-n-propylamine	U	5.00	"							
N-Nitrosodiphenylamine	U	5.00	"							
Pentachlorophenol	U	5.00	"							
Phenanthrene	U	5.00	"							
Phenol	U	5.00	"							
Pyrene	U	5.00	"							
1,2,4,5-Tetrachlorobenzene	U	5.00	"							
2,3,4,6-Tetrachlorophenol	U	5.00	"							
2,4,5-Trichlorophenol	U	5.00	"							
2,4,6-Trichlorophenol	U	5.00	"							
unknown (01)	8.96		"							T
unknown (02)	12.4		"							T
unknown (03)	3.61		"							T
Pentane, 2,3,4-trimethyl-	4.62		"							T
Cyclohexane, 1-methyl-2-propyl-	10.1		"							T
Surrogate: 2-Fluorophenol	71.3		"	100.00		71	21-110			
Surrogate: Phenol-d5	83.8		"	100.00		84	10-110			
Surrogate: Nitrobenzene-d5	37.6		"	50.000		75	35-114			
Surrogate: 2-Fluorobiphenyl	38.2		"	50.000		76	43-116			
Surrogate: 2,4,6-Tribromophenol	82.9		"	100.00		83	10-123			
Surrogate: Terphenyl-d14	44.0		"	50.000		88	33-141			



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Region 3 Environmental Science Center  
 Office of Analytical Services and Quality Assurance  
 701 Mapes Road  
 Fort Meade, Maryland 20755-5350



Site Name: **Dimock Residential Groundwater**

Project #: **DAS R33907**

**QC Data**  
**Semivolatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BA22504 - EPA 3520C SVOC**

LCS (BA22504-BS1)		Prepared: 01/25/12 12:43		Analyzed: 01/26/12 15:05					
Benzo(a)pyrene	3.94	5.00	ug/L	5.0000	79	30-150			J
Bis(2-chloroethyl)ether	4.10	5.00	"	5.0000	82	30-150			J
4-Chloroaniline	4.17	5.00	"	5.0000	83	30-150			J
4-Chloro-3-methylphenol	4.24	5.00	"	5.0000	85	26-103			J
2-Chlorophenol	4.28	5.00	"	5.0000	86	25-102			J
Diethyl phthalate	4.30	5.00	"	5.0000	86	30-150			J
2,4-Dinitrotoluene	3.89	5.00	"	5.0000	78	28-89			J
Hexachlorobenzene	4.09	5.00	"	5.0000	82	30-150			J
Hexachlorobutadiene	4.07	5.00	"	5.0000	81	30-150			J
Hexachloroethane	4.07	5.00	"	5.0000	81	30-150			J
Isophorone	4.19	5.00	"	5.0000	84	30-150			J
2-Methoxyethanol	13.5	5.00	"	23.160	58	30-150			
1-Methylnaphthalene	5.20	5.00	"	5.0000	104	30-150			
Naphthalene	3.96	5.00	"	5.0000	79	30-150			J
Nitrobenzene	4.10	5.00	"	5.0000	82	30-150			J
4-Nitrophenol	3.03	10.0	"	5.0000	61	11-114			J
N-Nitroso-di-n-propylamine	4.38	5.00	"	5.0000	88	41-126			J
N-Nitrosodiphenylamine	4.33	5.00	"	5.0000	87	30-150			J
Pentachlorophenol	1.18	5.00	"	5.0000	24	17-109			J
Phenol	4.22	5.00	"	5.0000	84	26-90			J
2,4,5-Trichlorophenol	4.30	5.00	"	5.0000	86	30-150			J
2,4,6-Trichlorophenol	4.24	5.00	"	5.0000	85	30-150			J
<i>Surrogate: 2-Fluorophenol</i>	<i>77.7</i>		<i>"</i>	<i>100.00</i>	<i>78</i>	<i>21-110</i>			
<i>Surrogate: Phenol-d5</i>	<i>89.6</i>		<i>"</i>	<i>100.00</i>	<i>90</i>	<i>10-110</i>			
<i>Surrogate: Nitrobenzene-d5</i>	<i>39.7</i>		<i>"</i>	<i>50.000</i>	<i>79</i>	<i>35-114</i>			
<i>Surrogate: 2-Fluorobiphenyl</i>	<i>39.9</i>		<i>"</i>	<i>50.000</i>	<i>80</i>	<i>43-116</i>			
<i>Surrogate: 2,4,6-Tribromophenol</i>	<i>89.2</i>		<i>"</i>	<i>100.00</i>	<i>89</i>	<i>10-123</i>			
<i>Surrogate: Terphenyl-d14</i>	<i>42.1</i>		<i>"</i>	<i>50.000</i>	<i>84</i>	<i>33-141</i>			



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Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22504 - EPA 3520C SVOC

Main data table with columns for analyte, result, limit, units, spike level, source result, %REC, and RPD. Includes sub-section 'LCS (BA22504-BS2)' and 'Surrogate' rows.



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Site Name: **Dimock Residential Groundwater**

Project #: **DAS R33907**

**QC Data**  
**Semivolatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BA22504 - EPA 3520C SVOC**

<b>Matrix Spike (BA22504-MS1)</b>	<b>Source: 1201013-01</b>			<b>Prepared: 01/25/12 12:43</b>		<b>Analyzed: 01/26/12 17:36</b>		
Benzo(a)pyrene	51.6	5.00	ug/L	60.000	0.00	86	30-150	
Bis(2-chloroethyl)ether	45.0	5.00	"	60.000	0.00	75	30-150	
4-Chloroaniline	55.8	5.00	"	60.000	0.00	93	30-150	
4-Chloro-3-methylphenol	56.2	5.00	"	60.000	0.00	94	26-103	
2-Chlorophenol	47.4	5.00	"	60.000	0.00	79	25-102	
Diethyl phthalate	53.5	5.00	"	60.000	0.00	89	30-150	
2,4-Dinitrotoluene	55.6	5.00	"	60.000	0.00	93	28-89	A
Hexachlorobenzene	46.6	5.00	"	60.000	0.00	78	30-150	
Hexachlorobutadiene	42.6	5.00	"	60.000	0.00	71	30-150	
Hexachloroethane	43.9	5.00	"	60.000	0.00	73	30-150	
Isophorone	52.1	5.00	"	60.000	0.00	87	30-150	
2-Methoxyethanol	35.7	5.00	"	57.900	0.00	62	30-150	
1-Methylnaphthalene	63.4	5.00	"	60.000	0.020	106	30-150	
Naphthalene	46.1	5.00	"	60.000	0.359	76	30-150	
Nitrobenzene	44.2	5.00	"	60.000	0.00	74	30-150	
4-Nitrophenol	61.4	10.0	"	60.000	0.00	102	11-114	
N-Nitroso-di-n-propylamine	52.0	5.00	"	60.000	0.00	87	41-126	
N-Nitrosodiphenylamine	50.7	5.00	"	60.000	0.00	85	30-150	
Pentachlorophenol	54.1	5.00	"	60.000	0.00	90	17-109	
Phenol	48.4	5.00	"	60.000	0.00	81	26-90	
2,4,5-Trichlorophenol	56.2	5.00	"	60.000	0.00	94	30-150	
2,4,6-Trichlorophenol	54.1	5.00	"	60.000	0.00	90	30-150	
<i>Surrogate: 2-Fluorophenol</i>	<i>72.3</i>		<i>"</i>	<i>100.00</i>		<i>72</i>	<i>21-110</i>	
<i>Surrogate: Phenol-d5</i>	<i>89.7</i>		<i>"</i>	<i>100.00</i>		<i>90</i>	<i>10-110</i>	
<i>Surrogate: Nitrobenzene-d5</i>	<i>39.3</i>		<i>"</i>	<i>50.000</i>		<i>79</i>	<i>35-114</i>	
<i>Surrogate: 2-Fluorobiphenyl</i>	<i>40.5</i>		<i>"</i>	<i>50.000</i>		<i>81</i>	<i>43-116</i>	
<i>Surrogate: 2,4,6-Tribromophenol</i>	<i>97.6</i>		<i>"</i>	<i>100.00</i>		<i>98</i>	<i>10-123</i>	
<i>Surrogate: Terphenyl-d14</i>	<i>44.4</i>		<i>"</i>	<i>50.000</i>		<i>89</i>	<i>33-141</i>	



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22603 - EPA 3520C SVOC

Blank (BA22603-BLK1)

Prepared: 01/26/12 12:30 Analyzed: 01/27/12 14:27

Main data table listing various chemical analytes (e.g., Acenaphthene, Benzaldehyde, etc.) with their corresponding results (U, 0.371, 0.077, 0.063, 1.90) and units (ug/L).



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Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22603 - EPA 3520C SVOC

Blank (BA22603-BLK1)

Prepared: 01/26/12 12:30 Analyzed: 01/27/12 14:27

Main data table listing various chemical compounds (e.g., Hexachlorobenzene, Nitrobenzene, Phenol) with their respective results, limits, and units.



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QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22603 - EPA 3520C SVOC

LCS (BA22603-BS1)

Prepared: 01/26/12 12:30 Analyzed: 01/27/12 15:18

Main data table listing various compounds like Benzo(a)pyrene, Bis(2-chloroethyl)ether, 4-Chloroaniline, etc., with their respective results and limits.



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QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22603 - EPA 3520C SVOC

LCS (BA22603-BS2)

Prepared: 01/26/12 14:28 Analyzed: 01/27/12 16:08

Main data table listing various compounds like Benzo(a)pyrene, Bis(2-chloroethyl)ether, etc., with their respective results, limits, and RPD values.



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Project #: **DAS R33907**

**QC Data**  
**Semivolatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BA22603 - EPA 3520C SVOC**

<b>Matrix Spike (BA22603-MS1)</b>	<b>Source: 1201013-05</b>			<b>Prepared: 01/26/12 12:30</b>		<b>Analyzed: 01/27/12 18:40</b>	
Benzo(a)pyrene	39.3	5.00	ug/L	60.000	0.00	66	30-150
Bis(2-chloroethyl)ether	38.2	5.00	"	60.000	0.00	64	30-150
4-Chloroaniline	42.9	5.00	"	60.000	0.00	71	30-150
4-Chloro-3-methylphenol	47.2	5.00	"	60.000	0.00	79	26-103
2-Chlorophenol	40.8	5.00	"	60.000	0.00	68	25-102
Diethyl phthalate	43.7	5.00	"	60.000	0.025	73	30-150
2,4-Dinitrotoluene	45.7	5.00	"	60.000	0.00	76	28-89
Hexachlorobenzene	37.3	5.00	"	60.000	0.00	62	30-150
Hexachlorobutadiene	37.0	5.00	"	60.000	0.00	62	30-150
Hexachloroethane	39.3	5.00	"	60.000	0.00	65	30-150
Isophorone	42.5	5.00	"	60.000	0.00	71	30-150
2-Methoxyethanol	27.1	5.00	"	57.900	0.00	47	30-150
1-Methylnaphthalene	51.1	5.00	"	60.000	0.00	85	30-150
Naphthalene	38.2	5.00	"	60.000	0.00	64	30-150
Nitrobenzene	37.2	5.00	"	60.000	0.00	62	30-150
4-Nitrophenol	53.3	10.0	"	60.000	0.00	89	11-114
N-Nitroso-di-n-propylamine	42.3	5.00	"	60.000	0.00	71	41-126
N-Nitrosodiphenylamine	38.9	5.00	"	60.000	0.00	65	30-150
Pentachlorophenol	44.1	5.00	"	60.000	0.00	73	17-109
Phenol	40.9	5.00	"	60.000	0.00	68	26-90
2,4,5-Trichlorophenol	44.9	5.00	"	60.000	0.00	75	30-150
2,4,6-Trichlorophenol	43.4	5.00	"	60.000	0.00	72	30-150
<i>Surrogate: 2-Fluorophenol</i>	<i>69.8</i>		<i>"</i>	<i>100.00</i>		<i>70</i>	<i>21-110</i>
<i>Surrogate: Phenol-d5</i>	<i>72.5</i>		<i>"</i>	<i>100.00</i>		<i>73</i>	<i>10-110</i>
<i>Surrogate: Nitrobenzene-d5</i>	<i>31.6</i>		<i>"</i>	<i>50.000</i>		<i>63</i>	<i>35-114</i>
<i>Surrogate: 2-Fluorobiphenyl</i>	<i>31.7</i>		<i>"</i>	<i>50.000</i>		<i>63</i>	<i>43-116</i>
<i>Surrogate: 2,4,6-Tribromophenol</i>	<i>73.7</i>		<i>"</i>	<i>100.00</i>		<i>74</i>	<i>10-123</i>
<i>Surrogate: Terphenyl-d14</i>	<i>32.2</i>		<i>"</i>	<i>50.000</i>		<i>64</i>	<i>33-141</i>



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701 Mapes Road
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22901 - EPA 3520C SVOC

Blank (BA22901-BLK1)

Prepared: 01/29/12 11:00 Analyzed: 01/30/12 19:39

Main data table listing various chemical analytes (e.g., Acenaphthene, Benzaldehyde, etc.) with their corresponding results (U, 0.318, 0.041, 0.718) and limits (5.00, 10.0).



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QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22901 - EPA 3520C SVOC

Blank (BA22901-BLK1)

Prepared: 01/29/12 11:00 Analyzed: 01/30/12 19:39

Main data table listing various compounds (Hexachlorobenzene, etc.) with columns for Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, and Notes.



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Site Name: Dimock Residential Groundwater

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QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22901 - EPA 3520C SVOC

Main data table with columns for analyte, result, limit, units, spike level, source result, %REC, %REC limits, RPD, RPD limit, and notes. Includes sub-section 'LCS (BA22901-BS1)' and 'Surrogate' rows.



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Site Name: **Dimock Residential Groundwater**

Project #: **DAS R33907**

**QC Data**  
**Semivolatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BA22901 - EPA 3520C SVOC**

**LCS (BA22901-BS2)**

Prepared: 01/29/12 11:00

Analyzed: 01/30/12 21:20

Benzo(a)pyrene	51.9	5.00	ug/L	60.000		86	30-150			
Bis(2-chloroethyl)ether	44.0	5.00	"	60.000		73	30-150			
4-Chloroaniline	54.7	5.00	"	60.000		91	30-150			
4-Chloro-3-methylphenol	54.5	5.00	"	60.000		91	26-103			
2-Chlorophenol	46.2	5.00	"	60.000		77	25-102			
Diethyl phthalate	52.1	5.00	"	60.000		87	30-150			
2,4-Dinitrotoluene	54.5	5.00	"	60.000		91	28-89			A
Hexachlorobenzene	46.5	5.00	"	60.000		78	30-150			
Hexachlorobutadiene	38.8	5.00	"	60.000		65	30-150			
Hexachloroethane	37.4	5.00	"	60.000		62	30-150			
Isophorone	50.6	5.00	"	60.000		84	30-150			
2-Methoxyethanol	35.0	5.00	"	57.900		60	30-150			
1-Methylnaphthalene	62.3	5.00	"	60.000		104	30-150			
Naphthalene	42.3	5.00	"	60.000		71	30-150			
Nitrobenzene	44.3	5.00	"	60.000		74	30-150			
4-Nitrophenol	60.2	10.0	"	60.000		100	11-114			
N-Nitroso-di-n-propylamine	50.5	5.00	"	60.000		84	41-126			
N-Nitrosodiphenylamine	50.8	5.00	"	60.000		85	30-150			
Pentachlorophenol	52.3	5.00	"	60.000		87	17-109			
Phenol	46.1	5.00	"	60.000		77	26-90			
2,4,5-Trichlorophenol	54.1	5.00	"	60.000		90	30-150			
2,4,6-Trichlorophenol	53.5	5.00	"	60.000		89	30-150			
<i>Surrogate: 2-Fluorophenol</i>	<i>72.0</i>		<i>ug/mL</i>	<i>100.00</i>		<i>72</i>	<i>21-110</i>			
<i>Surrogate: Phenol-d5</i>	<i>86.1</i>		<i>"</i>	<i>100.00</i>		<i>86</i>	<i>10-110</i>			
<i>Surrogate: Nitrobenzene-d5</i>	<i>39.7</i>		<i>"</i>	<i>50.000</i>		<i>79</i>	<i>35-114</i>			
<i>Surrogate: 2-Fluorobiphenyl</i>	<i>40.3</i>		<i>"</i>	<i>50.000</i>		<i>81</i>	<i>43-116</i>			
<i>Surrogate: 2,4,6-Tribromophenol</i>	<i>94.8</i>		<i>"</i>	<i>100.00</i>		<i>95</i>	<i>10-123</i>			
<i>Surrogate: Terphenyl-d14</i>	<i>44.5</i>		<i>"</i>	<i>50.000</i>		<i>89</i>	<i>33-141</i>			



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701 Mapes Road
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA22901 - EPA 3520C SVOC

Main data table with columns for Matrix Spike (BA22901-MS1), Source (1201013-15), Prepared (01/29/12 11:00), Analyzed (01/31/12 05:43), and various chemical analytes with their respective results and limits.



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701 Mapes Road
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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA23102 - EPA 3520C SVOC

Blank (BA23102-BLK1)

Prepared: 01/31/12 09:08 Analyzed: 02/01/12 16:59

Main data table listing various chemical analytes (e.g., Acenaphthene, Benzaldehyde, etc.) with their corresponding results (U, 1.31, 0.052, 0.367, 0.485) and units (ug/L).



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA23102 - EPA 3520C SVOC

Blank (BA23102-BLK1)

Prepared: 01/31/12 09:08 Analyzed: 02/01/12 16:59

Main data table listing various compounds (e.g., Hexachlorobenzene, Hexachlorobutadiene, etc.) with their respective results, limits, and units.

Summary table for surrogate compounds: Surrogate: 2-Fluorophenol, Surrogate: Phenol-d5, Surrogate: Nitrobenzene-d5



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Site Name: Dimock Residential Groundwater

Project #: DAS R33907

QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA23102 - EPA 3520C SVOC

Blank (BA23102-BLK1)

Prepared: 01/31/12 09:08 Analyzed: 02/01/12 16:59

Table with 6 columns: Surrogate, Result, Units, Spike Level, %REC, %REC Limits

LCS (BA23102-BS1)

Prepared: 01/31/12 09:08 Analyzed: 02/01/12 17:50

Main table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes



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Site Name: Dimock Residential Groundwater

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QC Data
Semivolatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BA23102 - EPA 3520C SVOC

LCS (BA23102-BS2)

Prepared: 01/31/12 09:08 Analyzed: 02/01/12 18:41

Main data table listing various compounds like Benzo(a)pyrene, Bis(2-chloroethyl)ether, etc., with their respective results, limits, and RPD values.



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Project #: DAS R33907

QC Data  
 Semivolatile Organic Compounds

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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Batch BA23102 - EPA 3520C SVOC

Matrix Spike (BA23102-MS1)	Source: 1201013-34			Prepared: 01/31/12 09:08	Analyzed: 02/02/12 02:18	
Benzo(a)pyrene	44.7	5.00	ug/L	60.000	0.00	74 30-150
Bis(2-chloroethyl)ether	41.9	5.00	"	60.000	0.00	70 30-150
4-Chloroaniline	48.4	5.00	"	60.000	0.00	81 30-150
4-Chloro-3-methylphenol	48.7	5.00	"	60.000	0.00	81 26-103
2-Chlorophenol	44.6	5.00	"	60.000	0.00	74 25-102
Diethyl phthalate	45.2	5.00	"	60.000	0.273	75 30-150
2,4-Dinitrotoluene	47.8	5.00	"	60.000	0.00	80 28-89
Hexachlorobenzene	42.0	5.00	"	60.000	0.00	70 30-150
Hexachlorobutadiene	38.9	5.00	"	60.000	0.00	65 30-150
Hexachloroethane	41.0	5.00	"	60.000	0.00	68 30-150
Isophorone	43.2	5.00	"	60.000	0.00	72 30-150
2-Methoxyethanol	37.2	5.00	"	57.900	0.00	64 30-150
1-Methylnaphthalene	53.9	5.00	"	60.000	0.00	90 30-150
Naphthalene	41.7	5.00	"	60.000	0.00	70 30-150
Nitrobenzene	41.4	5.00	"	60.000	0.00	69 30-150
4-Nitrophenol	53.1	10.0	"	60.000	0.00	89 11-114
N-Nitroso-di-n-propylamine	46.3	5.00	"	60.000	0.00	77 41-126
N-Nitrosodiphenylamine	45.1	5.00	"	60.000	0.00	75 30-150
Pentachlorophenol	43.6	5.00	"	60.000	0.00	73 17-109
Phenol	45.4	5.00	"	60.000	0.00	76 26-90
2,4,5-Trichlorophenol	49.2	5.00	"	60.000	0.00	82 30-150
2,4,6-Trichlorophenol	50.4	5.00	"	60.000	0.00	84 30-150
Surrogate: 2-Fluorophenol	75.4		"	100.00		75 21-110
Surrogate: Phenol-d5	81.5		"	100.00		82 10-110
Surrogate: Nitrobenzene-d5	35.4		"	50.000		71 35-114
Surrogate: 2-Fluorobiphenyl	36.1		"	50.000		72 43-116
Surrogate: 2,4,6-Tribromophenol	87.0		"	100.00		87 10-123
Surrogate: Terphenyl-d14	41.5		"	50.000		83 33-141



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Blank (BB20301-BLK1)

Prepared & Analyzed: 01/27/12 12:14

Table listing various chemical compounds (e.g., Acetone, Benzene, Bromobenzene) with their corresponding results (U) and quantitation limits (e.g., 2.0 ug/L).



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**QC Data**  
**Volatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BB20301 - VOC Purge and Trap**

**Blank (BB20301-BLK1)**

Prepared & Analyzed: 01/27/12 12:14

Hexachlorobutadiene	U	0.5	ug/L							
2-Hexanone	U	2.0	"							
Isopropylbenzene	U	0.5	"							
p-Isopropyltoluene	U	0.5	"							
Methyl Acetate	U	0.5	"							
Methylcyclohexane	U	0.5	"							
Methyl-tert-butyl ether	U	0.5	"							
Methylene Chloride	U	0.5	"							
4-Methyl-2-pentanone	U	2.0	"							
Naphthalene	U	0.5	"							
n-Propylbenzene	U	0.5	"							
Styrene	U	1.0	"							
1,1,2,2-Tetrachloroethane	U	0.5	"							
1,1,1,2-Tetrachloroethane	U	0.5	"							
Tetrachloroethene	U	0.5	"							
Toluene	U	0.5	"							
1,2,3-Trichlorobenzene	U	0.5	"							
1,2,4-Trichlorobenzene	U	0.5	"							
1,1,1-Trichloroethane	U	0.5	"							
1,1,2-Trichloroethane	U	0.5	"							
Trichloroethene	U	0.5	"							
Trichlorofluoromethane	U	0.5	"							
1,2,3-Trichloropropane	U	0.5	"							
1,2,4-Trimethylbenzene	U	0.5	"							
1,3,5-Trimethylbenzene	U	0.5	"							
Vinyl acetate	U	0.5	"							
Vinyl chloride	U	0.5	"							
m-Xylene/p-Xylene	U	1.0	"							
o-Xylene	U	1.0	"							
<i>Surrogate: 4-Bromofluorobenzene</i>	<i>3.790</i>	<i>"</i>	<i>"</i>	<i>4.0000</i>	<i>95</i>	<i>86-115</i>				
<i>Surrogate: 1,2-Dichloroethane-d4</i>	<i>4.260</i>	<i>"</i>	<i>"</i>	<i>4.0000</i>	<i>106</i>	<i>76-114</i>				
<i>Surrogate: Toluene-d8</i>	<i>3.920</i>	<i>"</i>	<i>"</i>	<i>4.0000</i>	<i>98</i>	<i>88-110</i>				



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Blank (BB20301-BLK2)

Prepared & Analyzed: 01/30/12 13:05

Table listing various chemical compounds (e.g., Acetone, Benzene, Bromobenzene) with their corresponding results (U) and quantitation limits (e.g., 2.0 ug/L).



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Blank (BB20301-BLK2)

Prepared & Analyzed: 01/30/12 13:05

Main data table listing various chemical compounds (e.g., Hexachlorobutadiene, 2-Hexanone, Isopropylbenzene) with their respective results (U), quantitation limits (0.5, 2.0, 1.0 ug/L), and surrogate data (e.g., 4-Bromofluorobenzene, 1,2-Dichloroethane-d4, Toluene-d8).



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Blank (BB20301-BLK3)

Prepared & Analyzed: 01/31/12 13:31

Table listing various chemical compounds (e.g., Acetone, Benzene, Bromobenzene) with their corresponding results (U) and quantitation limits (e.g., 2.0 ug/L).



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**QC Data**  
**Volatile Organic Compounds**

Analyte	Result	Quantitation Limit	Units	Spike Level	Source Result	%REC	%REC Limits	RPD	RPD Limit	Notes
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**Batch BB20301 - VOC Purge and Trap**

**Blank (BB20301-BLK3)**

Prepared & Analyzed: 01/31/12 13:31

Hexachlorobutadiene	U	0.5	ug/L							
2-Hexanone	U	2.0	"							
Isopropylbenzene	U	0.5	"							
p-Isopropyltoluene	U	0.5	"							
Methyl Acetate	U	0.5	"							
Methylcyclohexane	U	0.5	"							
Methyl-tert-butyl ether	U	0.5	"							
Methylene Chloride	U	0.5	"							
4-Methyl-2-pentanone	U	2.0	"							
Naphthalene	U	0.5	"							
n-Propylbenzene	U	0.5	"							
Styrene	U	1.0	"							
1,1,2,2-Tetrachloroethane	U	0.5	"							
1,1,1,2-Tetrachloroethane	U	0.5	"							
Tetrachloroethene	U	0.5	"							
Toluene	U	0.5	"							
1,2,3-Trichlorobenzene	U	0.5	"							
1,2,4-Trichlorobenzene	U	0.5	"							
1,1,1-Trichloroethane	U	0.5	"							
1,1,2-Trichloroethane	U	0.5	"							
Trichloroethene	U	0.5	"							
Trichlorofluoromethane	U	0.5	"							
1,2,3-Trichloropropane	U	0.5	"							
1,2,4-Trimethylbenzene	U	0.5	"							
1,3,5-Trimethylbenzene	U	0.5	"							
Vinyl acetate	U	0.5	"							
Vinyl chloride	U	0.5	"							
m-Xylene/p-Xylene	U	1.0	"							
o-Xylene	U	1.0	"							
<i>Surrogate: 4-Bromofluorobenzene</i>	<i>3.780</i>		<i>"</i>	<i>4.0000</i>		<i>94</i>	<i>86-115</i>			
<i>Surrogate: 1,2-Dichloroethane-d4</i>	<i>4.330</i>		<i>"</i>	<i>4.0000</i>		<i>108</i>	<i>76-114</i>			
<i>Surrogate: Toluene-d8</i>	<i>3.600</i>		<i>"</i>	<i>4.0000</i>		<i>90</i>	<i>88-110</i>			



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Blank (BB20301-BLK4)

Prepared & Analyzed: 02/02/12 11:56

Table listing various chemical compounds (e.g., Acetone, Benzene, Bromobenzene) with their corresponding results (U) and quantitation limits (e.g., 2.0 ug/L, 0.5).



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Blank (BB20301-BLK4)

Prepared & Analyzed: 02/02/12 11:56

Main data table listing various chemical compounds (e.g., Hexachlorobutadiene, 2-Hexanone, Isopropylbenzene) with their respective results (U), quantitation limits (0.5, 2.0, 1.0 ug/L), and surrogate data (4-Bromofluorobenzene, 1,2-Dichloroethane-d4, Toluene-d8).



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

LCS (BB20301-BS1)

Prepared & Analyzed: 01/24/12 19:56

Main data table listing various analytes (e.g., Acetone, Benzene, Bromobenzene) with their respective results, limits, units, spike levels, source results, %REC, RPD, and notes.



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

LCS (BB20301-BS1)

Prepared & Analyzed: 01/24/12 19:56

Main data table listing various VOCs (Hexachlorobutadiene, 2-Hexanone, etc.) with columns for Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, and Notes.



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Main data table with columns for analyte, result, limit, units, spike level, source result, %REC, %REC limits, RPD, RPD limit, and notes. Includes sub-headers for Matrix Spike and Source information.



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Matrix Spike (BB20301-MS1) Source: 1201013-14 Prepared & Analyzed: 02/02/12 12:50. Table listing various chemical compounds and their results.

Matrix Spike (BB20301-MS2) Source: 1201013-33 Prepared & Analyzed: 02/02/12 13:17. Table listing various chemical compounds and their results.



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Main data table with columns: Matrix Spike (BB20301-MS2), Source: 1201013-33, Prepared & Analyzed: 02/02/12 13:17, and various chemical analytes with their respective results and limits.



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QC Data
Volatile Organic Compounds

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes

Batch BB20301 - VOC Purge and Trap

Table with 11 columns: Analyte, Result, Quantitation Limit, Units, Spike Level, Source Result, %REC, %REC Limits, RPD, RPD Limit, Notes. Includes sub-headers for Matrix Spike and Source information.



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**Notes and Definitions**

- UJ The analyte was not detected at or above the quantitation limit. The quantitation limit is an estimate.
- T Tentatively Identified Compound. Identified as a result of a library search using the EPA/NIST Mass Spectral Library. Standards were not used to verify the identity and quantity of the compound. The reported value is an estimate.
- K The identification of the analyte is acceptable; the reported value may be biased high. The actual value is expected to be less than the reported value. Reported value is an estimate.
- J The identification of the analyte is acceptable; the reported value is an estimate.
- B Not detected substantially above (10 times) the level reported in the laboratory or field blanks (including field, trip, rinsate, and equipment blanks).
- A Quality control value is outside acceptance limits.
- NR Not Reported
- RPD Relative Percent Difference
- U Analyte included in the analysis, but not detected at or above the quantitation limit.

Quantitation Limit: The lowest concentration of an analyte that can be reliably measured within specified limits of precision and accuracy for a specific laboratory analytical method and that takes into account analytical adjustments made during sample preparation and analysis.

SOLID SAMPLE RESULTS - REPORTING PROTOCOL: Solid samples where % Solids (percent dry wt at 105 degrees C) has been performed, are analyzed wet and converted to a dry weight result for reporting purposes. This is routine for organics and most inorganic analyses. When metals and mercury analyses are requested, solid samples are routinely analyzed and reported on a dry weight basis. Solid samples for metals/mercury are prepared for analysis by an initial drying at 60 degree C and homogenization before digestion. Oil-type samples will be analyzed and reported on a wet weight basis for all analyses because of the nature of the sample. Any exceptions to the protocol will be noted with a qualifier